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(11) EP 1 284 298 A2

(12)

EUROPEAN PATENT APPLICATION

(43) Date of publication: 19.02,2003 Bulletin 2003/08

(51) Int Cl.7: C12Q 1/68, A01K 67/027

(21) Application number: 02255229.3

(22) Date of filing: 26.07.2002

(84) Designated Contracting States:

AT BE BG CH CY CZ DE DK EE ES FI FR GB GR
IE IT LI LU MC NL PT SE SK TR
Designated Extension States:

AL LT LV MK RO SI

(30) Priority: 27.07.2001 GB 0118354 07.02.2002 GB 0202883

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(54) Identification and use of molecules implicated in pain

(57) The invention relates to the use of:

(a) an isolated gene sequence that is down-regulated in the spinal cord of a mammal in response to mechanistically distinct first and second models of neuropathic or central sensitization pain;

- (b) an isolated gene sequence comprising a nucleic acid sequence of any of Tables I to VI;
- (c) an isolated gene sequence having at least 80% sequence identity with a nucleic acid sequence of any of Tables I to VI;
- (d) an isolated nucleic acid sequence that is hybridizable to any of the gene sequences according to
 (a), (b) or (c) under stringent hybridisation conditions;
- (e) a recombinant vector comprising a gene sequence or nucleic acid sequence according to any one of (a) to (d);
- (f) a host cell containing the vector according to (e);
 (g) a non-human animal having in its genome an introduced gene sequence or nucleic acid sequence or a removed or down-regulated gene sequence or nucleic acid sequence according to any one of (a) to (d);
- (h) an isolated polypeptide comprising an amino acid sequence at least 90% identical to an amino acid sequence encoded by a nucleotide sequence ac-

cording to any one of (a) to (d), or a variant polypeptide thereof with sequential amino acid deletions from the C terminus and/or the N-terminus;

- (i) an isolated polypeptide encoded by a nucleotide sequence according to any one of (a) to (d); or
- (j) an isolated antibody that binds specifically to a polypeptide according to (h) or (i);

in the screening of compounds for the treatment of pain, or for the diagnosis of pain.

The invention also relates to the use of naturally occurring compounds such as peptide ligands of the expression products of the above gene sequences and their associated signal transduction pathways for use in the treatment of pain.

Description

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FIELD OF THE INVENTION

[0001] The present invention relates to nucleic acids, their expression products and pathways involved in paln, and their use in screening for molecules that can alleviate pain. The invention further relates to methods for the assay and diagnosis of pain in patients.

BACKGROUND TO THE INVENTION

[0002] Pain is currently classified into four general types. Post-operative acute pain can be successfully treated with existing pain medications of e.g. the opioid and non-steroidal anti-inflammatory (NSAID) types, and is usually short-term and self-limiting. A second type of pain, present e.g. in cancer and arthritis, is also responsive to medication initially with a NSAID and in its later stages with opioids. Neuropathic pain arises from damage to the central or peripheral nerve systems, and is more effectively treated with antidepressants or anticonvulsants. A fourth type of pain called central sensitization results from changes in the central nervous system as a result of chronic pain, these changes often being irreversible and difficult to treat. Nerve pain from shingles or diabetes falls into this and the neuropathic category. Changes occur where pain is at first poorly controlled and gradually progress to the point where a person is sensitive to stimuli which would not normally cause pain, for example a light touch. People with pain of this kind often describe a widening of the pain area to include areas which had originally not been injured or which were thought not to be involved in pain. This classification is, however based on clinical symptoms rather than on the underlying pain mechanisms.

[0003] Opiates such as morphine belong to a traditional class of pain-relieving compounds that are now recognized as binding to opiate receptors. Naturally occurring polypeptides have also been found to have opiate-like effects on the central nervous system, and these include β-endorphin, met-enkephalin and leu-enkephalin.

[0004] Salicin was isolated at the beginning of the 19th century, and from that discovery a number of NSAIDs such as aspirin, paracetamol, ibuprofen, flurbiprofen and naproxen were developed. NSAIDs are by far the most widely used pain-relieving compounds, but can exhibit side effects, in particular irritation of the GI tract that can lead to the formation of ulcers, gastrointestinal bleeding and anemia.

[0005] Interest in the neurobiology of pain is developing: see a colloquium sponsored by the US National Academy of Sciences in December 1998 concerning the neurobiology of pain and reviewed in *The Scientist* 13[1], 12, 1999. Many pain mechanisms were discussed including the role of the capsaicin receptors in pain, (M.J. Caterina *et al.*, *Nature*, 389, 816-824, 1997). Large dosages of capsaicin were reported to disable that receptor, (W.R. Robbins *et al.*, *Anesthesia and Analgesia*, 86, 579-583, 1998). Additionally, a tetrodotoxin-resistant sodium channel found in small diameter pain-sensing neurons (PN3) was discussed (A.N. Akopian *et al.*, *Nature*, 379, 257-262, 1986) and L. Sangameswaran *et al.*, *Journal of Biological Chemistry*, 271, 953-956, 1996). Its involvement in transmission and sensitization to pain signals has been reported, (S.D. Novakovic *et al.*, *Journal of Neuroscience*, 18, 2174-2187, 1998). A further tetrodotoxin-resistant sodium channel has been reported (S. Tate *et al.*, *Nature Neuroscience*, 1, 653-655 1998). [0006] Second messenger systems have also been shown to be important since knockout-mice lacking protein kinase C (PKC) γ were reported to respond to acute pain e.g. from a hot surface, but not to respond to neuropathic pain when their spinal nerves are injured (Malmberg *et al.*, *Science*, 278, 279-283 (1997).

[0007] Present methods for identifying novel compounds that relieve pain of one or more of the types indicated above suffer from the defect that they are dependent either on the relatively limited number of receptors known to be involved in pain or on the empirical identification of new receptors which is an uncertain process. In relation to known receptors, for example the opioid receptor, research directed to improved compounds offers the possibility of screenining compounds that have a better therapeutic ratio and fewer side effects. This does not lead naturally to compounds for different pain receptors that have new modes of action and new and qualitatively different benefits. Even when newly identified additional receptors are taken into account, known receptors revolve around tens of gene products. However, there are between 30,000 and 40,000 genes in the genome of an animal and more of them are concerned with nervous system function than with peripheral function. We therefore concluded that a large number of receptors and pathways are important to the transduction of pain, but up to now have remained unknown.

SUMMARY OF THE INVENTION

[0008] It is an object of the invention to provide sequences of genetic material for which no role in pain has previously been disclosed, and which are useful, for example, in:

· identifying metabolic pathways for the transduction of pain

- identifying from said metabolic pathways compounds having utility in the diagnosis or treatment of pain
- producing proteins and polypeptides with a role in the transduction of pain;
- producing genetically modified non-human animals that are useful in the screening of compounds having utility in the treatment or diagnosis of pain.
- Identifying ligand molecules for receptors involved in said metabolic pathways and having utility in the treatment of pain.

[0009] It is yet a further object of the invention to provide research tools, for example non-human animals and microorganisms, that can be used in screening compounds for pharmacological activity, especially pain-reducing activity.

[0010] The present invention is based on sequences that are down-regulated in two models of chronic pain, namely streptozocin-induced diabetes and chronic constrictive injury (CCI) to a nerve leading to the spine, for example the sciatic nerve.

[0011] In one aspect, the invention relates to the use in the screening of compounds that are effective in the treatment of pain, or in the diagnosis of pain, of:

- (a) an isolated gene sequence that is down regulated in the spinal cord of a mammal in response to first and second models of pain, for example in response to streptozocin-induced diabetes and in response to a chronic constrictive injury to a nerve leading into the spine;
- (b) an isolated gene sequence having at least 80% sequence identity with any of the nucleic acid sequences of Tables I VI, preferably 85% sequence identity, more preferably 90%, increasingly preferably 95%, most preferably 99%.
- (c) an isolated nucleic acid sequence comprising a sequence that is hybridizable to any of the gene sequences according to (a) or (b) under stringent hybridisation conditions;
- (d) a recombinant vector comprising any of the gene sequences according to (a) to (c);
- (e) a host cell containing the vector according to (d);
- (f) a non-human animal, for use in the screening of compounds that are effective in the treatment of pain, or in the diagnosis of pain, having in its genome an introduced gene sequence or a removed or down-regulated nucleotide sequence, said sequence becoming down-regulated in the spinal cord of a mammal in response to first and second models of pain, particularly neuropathic or sensitisation pain, for example in response to streptozocin-induced diabetes and in response to a chronic constrictive injury to a nerve leading into the spine;
- (g) an isolated polypeptide containing an amino acid sequence at least 90% identical to an amino acid sequence encoded by a nucleotide sequence according to any one of (a) to (d), or a variant thereof with sequential amino acid deletions from the C terminus and/or the N-terminus; or
- (h) an isolated antibody that binds specifically to the isolated polypeptide according to (g).

[0012] The invention further provides a compound that is useful in the treatment or diagnosis of pain and that modulates the action of an expression product of a gene sequence that becomes down-regulated in the spinal cord of a mammal in response to first and second models of pain, for example being down-regulated both in response to streptozocin induced diabetes and in response to chronic constrictive injury to a nerve leading into the spine.

[0013] The invention also relates to the use of naturally occurring compounds such as peptide ligands of the expression products of the above gene sequences and their associated signal transduction pathways for the treatment of pain.

DESCRIPTION OF PREFERRED EMBODIMENTS

45 DEFINITIONS

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[0014] Within the context of the present invention:

- "Comprising" means consisting of or including. Thus nucleic acid comprising a defined sequence includes nucleic
 acid that may contain a full-length gene or full-length cDNA. The gene may include any of the naturally occurring
 regulatory sequence(s), such as a transcription and translation start site, a promoter, a TATA box in the case of
 eukaryotes, and transcriptional and translational stop sites. Further, nucleic acid comprising a cDNA or gene may
 include any appropriate regulatory sequences for the efficient expression thereof in vitro.
- "Isolated" requires that the material be removed from its original environment (e.g. the natural environment if it is naturally occurring). For example, a naturally occurring polynucleotide or a peptide present in a living animal is not isolated, but the same polynucleotide or peptide, separated from some or all of the coexisting materials in the natural system, is isolated. Such polynucleotide can be part of a vector and/or such polynucleotide or peptide can

be part of a composition, and still be isolated in that the vector or composition is not a part of its natural environment.

- "Mechanistically distinct" in relation to pain models implies that the pain is induced by mechanisms that differ in kind rather than being variants of a similar pain model. Thus diabetic pain and chronic constrictive pain models are mechanistically distinct, whereas spinal nerve ligation models and sciatic nerve ligation models which both work by ligation are not.
- "Purified" does not require absolute purity; instead it is intended as a relative definition. Purification of starting
 materials or natural materials from their native environment to at least one order of magnitude, preferably two or
 three orders, and more preferably four or five orders of magnitude is expressly contemplated.
- "Nucleic acid sequence" or "gene sequence" means a sequence of nucleotides or any variant or homologue thereof, or truncated or extended sequence thereof, and is preferably indicated by a Genebank accession number. Also within the scope of the present invention are down-regulated nucleic acid sequences which encode expression products which are components of signaling pathways. This invention also includes any variant or homologue or truncated or extended sequence of the down-regulated nucleotide sequence. Also within the scope of the present invention, the term "nucleic acid(s) product", or "expression product" or "gene product" or a combination of these terms refers without being biased, to any, protein(s), polypeptide(s), peptide(s) or fragment(s) encoded by the down-regulated nucleotide sequence.
- "Operably linked" refers to a linkage of polynucleotide elements in a functional relationship. For instance, a promoter or an enhancer is operably linked to a coding sequence if it regulates the transcription of the coding sequence. In particular, two DNA molecules (such as a polynucleotide containing a promoter region, and a polynucleotide encoding a desired polypeptide) are said to be "operably linked" if the nature of the linkage between the two polynucleotides does not (1) result in the introduction of a frame-shift mutation and (2) interfere with the ability of the polynucleotide containing the promoter to direct the transcription of the coding polynucleotide.

"Gene product" refers to polypeptide - which is interchangeable with the term protein - which is encoded by a nucleotide sequence and includes single-chain polypeptide molecules as well as multiple-polypeptide complexes where individual constituent polypeptides are linked by covalent or non-covalent means. Polypeptides of the present invention may be produced by synthetic means (e.g. as described by Geysen *et al.*, 1996) or by recombinant means.

The terms "variant", "homologue", "fragment", "analogue" or "derivative" in relation to the amino acid sequence for the polypeptide of the present invention includes any substitution of, variation of, modification of, replacement of, deletion of or addition of one (or more) amino acid from or to the sequence providing the resultant polypeptide has the native gene product activity. In particular, the term "homologue" covers homology with respect to structure and/or function. With respect to sequence homology, there is at least 90%, more preferably at least 95% homology to an amino acid sequence encoded by the relevant nucleotide sequence shown in Tables I - VI, preferably there is at least 98% homology.

Typically, for the variant, homologue or fragment of the present invention, the types of amino acid substitutions that could be made should maintain the hydrophobicity/hydrophilicity of the amino acid sequence. Amino acid substitutions may include the use of non-naturally occurring amino acid analogues.

In addition, or in the alternative, the protein itself could be produced using chemical methods to synthesize a polypeptide, in whole or in part. For example, peptides can be synthesized by solid phase techniques, cleaved from the resin, and purified by preparative high performance liquid chromatography (e.g. Creighton (1983) Proteins Structures and Molecular Principles, WH Freeman and Co., New York, NY, USA). The composition of the synthetic peptides may be confirmed by amino acid analysis or sequencing (e.g. the Edman degradation procedure).

Direct peptide synthesis can be performed using various solid-phase techniques (Roberge JY et al Science Vol 269 1995 202-204) and automated synthesis may be achieved, for example, using the ABI 431 A Peptide Synthesizer (Perkin Elmer) in accordance with the instructions provided by the manufacturer. Additionally, the amino acid sequence of a gene product, or any part thereof, may be altered during direct synthesis and/or combined using chemical methods with a sequence from other subunits, or any part thereof, to produce a variant polypeptide.

In another embodiment of the invention, a gene product natural, modified or recombinant amino acid sequence may be ligated to a heterologous sequence to encode a fusion protein. For example, for screening of libraries for compounds and peptide agonists and antagonists of gene product activity, it may be useful to encode a chimeric gene product expressing a heterologous epitope that is recognised by a commercially available antibody. A fusion protein may also be engineered to contain a cleavage site located between a gene product sequence and the heterologous protein sequence, so that the gene product may be cleaved and purified away from the heterologous moiety.

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The gene product may also be expressed as a recombinant protein with one or more additional polypeptide domains added to facilitate protein purification. Such purification facilitating domains include, but are not limited to, metal chelating peptides such as histidine-tryptophan modules that allow purification on immobilised metals (Porath J, Protein Expr Purif Vol 3 1992 p263-281), protein A domains that allow purification on immobilised immunoglobulin, and the domain utilised in the FLAGS extension/affinity purification system (Immunex Corp, Seattle, WA, USA). The inclusion of a cleavable linker sequence such as Factor XA or enterokinase (Invitrogen, San Diego, CA, USA) between the purification domain and the gene product is useful to facilitate purification.

- "Pain" includes chronic pain, neuropathic pain, pain arising from central sensitisation, and in particular diabetic pain.
- "Stringent hybridization conditions" is a recognized term in the art and for a given nucleic acid sequence refers to those conditions which permit hybridization of that sequence to its complementary sequence and closely homologous sequences. Conditions of high stringency may be illustrated in relation to filter-bound DNA as for example 2X SSC, 65°C (where SSC = 0.15M sodium chloride, 0.015M sodium citrate, pH 7.2), or as 0.5M NaHPO₄, 7% sodium dodecyl sulfate (SDS), 1mM EDTA, at 65°C, and washing in 0.1xSCC/0.1% SDS at 68°C (Ausubel F.M. et al., eds, 1989, Current Protocols in Molecular Biology, Vol. I, Green Publishing Associates, Inc., and John Wiley & Sons Inc., New York, at p. 2. 10.3). Hybridization conditions can be rendered highly stringent by raising the temperature and/or by the addition of increasing amounts of formamide, to destabilize the hybrid duplex of non-homologous nucleic acid sequence relative to homologous and closely homologous nucleic acid sequences. Thus, particular hybridisation conditions can be readily manipulated, and will generally be chosen depending on the desired results.
- "Variants or homologues" include (a) sequence variations of naturally existing gene(s) resulting from polymorphism (s), mutation(s), or other alteration(s) as compared to the above identified sequences, and which do not deprive the encoded protein of function (b) recombinant DNA molecules, such as cDNA molecules encoding genes indicated by the relevant Genebank accession numbers and (c) any sequence that hybridizes with the above nucleic acids under stringent conditions and encodes a functional protein or fragment thereof.

IDENTIFIED SEQUENCES

[0015] The inventors have identified nucleotide sequences that give rise to expression products listed in Tables I - VI, that become differentially expressed in the spinal cord in response to two distinct chronic pain stimuli, for example neuropathic and/or central sensitization pain stimuli, and that are believed to be involved in the transduction of pain. In Tables I - VI, * denotes more preferred nucleic acid sequences and ** denotes most preferred nucleic acid sequences. These nucleic acid sequences have not previously been implicated in the transduction of pain.

[0016] The validity of the present experimental procedure was confirmed by the fact that nucleotide sequences were obtained as a result of the investigation whose function in the transduction of pain has been previously confirmed and established. These nucleic acid sequences are not part of this invention. Any of the nucleic acid sequences and expression products can be used to develop screening technologies for the identification of novel molecules for the prevention or treatment of pain. These screening technologies could also be used to ascribe new pain therapeutic indications to molecules that have not previously been identified as being useful for the prevention or treatment of pain. Furthermore, the said nucleic acid sequences can be used as diagnostic tools and for the development of diagnostic tools.

Table I -

Sequences whos	e expression produc	cts are kinases			
Expression productor name (Reference)	Rat Accession Number	Mouse accession number	Human Accession Number	Reaction	Assay
Pyruvate kinase, M1 and M2 subunits (Refs:1 -2)	M24359 (SEQ ID No's 1-3)	X97047	X56494	Tissue-specific promoter; Carbohydrate kinase	Kinase

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Table II -

Expression Product or Name (Reference)	Rat Accession Number	Mouse accession Number	Human Accession Number	Reaction	Assay
Dopamine receptor D.sub. 1** (Ref : 3)	I58000 (SEQ ID NO 4)			Dopamine receptor	Receptor
Putative GABA-B1a receptor** (Ref:		AF114168 (SEQ ID No's 5-6)		GABA-B Receptor	Receptor

Table III -

20	Sequences who	se expression pro	ducts are transpo	rters		
	Expression Product or Name	Rat Accession Number	Mouse Accession Number	Human Accession Number	Reaction	Assay
25 30	Differentiation- associated Na- dependent inorganic phosphate cotransporter (Ref: N/A)	AF271235(SEQ ID NO's 7-8)			Trans-membrane phosphate transport	Transporter
35	Putative vacuolar assembly protein VSP41 gene (Ref : N/A)	U87309		AAM47563. 1 (SEQ ID No 9)	Vacuolar assembly and traffic	Transporter

Table IV -

Sequences whos	se expression prod	lucts are G-prote	in coupled recepto	or proteins	
Expression product or Name	Rat Accession Number	Mouse Accession Number	Human Accession Number	Function	Assay
Git1 (G-protein- coupled receptor kinase- interactor 1; GPCR kinase- associated ADP- ribosylation factor) (Ref: 5)	AF085693 (SEQ ID NO 10-11)			Regulation of activity of ARF6 in phosphatidylinIsitol 3-kInase signalling pathways; β2 adrenergic receptor regulation	Ligand binding

Table V -

Sequences whose	e expression produ	cts are DNA-bind	ing proteins		
Expression Product or Name	Rat Accession Number	Mouse Accession Number	Human Accession Number	Reaction	Assay
Putative histone H3.3A (Ref: 6)		X91866	M11354(SEQ ID NO 12-13)	DNA binding	DNA binding

Table VI -

	Sequences whose	expression produ	ıcts are other er	ızymes		
15	Expression product or Name	Rat Accession Number	Mouse Accession Number	Human Accession Number	Reaction	Assay
20	3-Hydroxy 3-methylglutaryl coenzyme A synthase, cytosolic * (Ref: 7)	X52625 (SEQ ID No's 14-15)			Cholesterol biosynthesis	Ligase
25	Acyl-CoA synthetase II, brain (Ref : N/A)	D360666 (N/A)			Fatty acid metabolism	Ligase
30	Farnesyl diphosphate synthase ** (Ref : 8)	M34477 (Seq ID No's 16-17)			Isoprene biosynthesis	Ligase
	Bendless protein (Ref : N /A)	AB032739 (Seq ID No's 18-19)		E12457	Protein degradation	Ligase
35	fatty acid synthase (Ref: 9)	X62888 (SEQ ID No's 20-21)			Fatty acid synthesis	Ligase
40	Glutamine synthetase (EC 6.3.1.2) ** (Ref: 10)	M91652 (SEQ ID NO's 22-23)			Amino acid metabolism	Ligase
45	Putative seryl- tRNA synthetase (Ref No : 11)			X91257 (SEQID NO's 24-25)	Ligase	Ligase
45	Enolase, alpha alpha, non- neuronal (NNE) (REF NO 12)	X02610	X52379	M14328 (SEQ ID NO's 26-27)	Glycolysis	Lyase
50	Aldose reductase, lens (AREC 11.1.21) (REF: 13)	X05884 (SEQ ID NO's 28-29)			Reduces carbonyl	Oxidoreductase
55	Cytochrome-c oxidase I, mitochondrial (Ref : 14)	S79304 (SEQ ID NO's 30-31)			Mitochrondrial energy metabloism	Oxidoreductase

Table VI - (continued)

	expression produ			B - 11 - 1	T A
Expression product or Name	Rat Accession Number	Mouse Accession Number	Human Accession Number	Reaction	Assay
Lactate dehydrogenase-B (LDH-B) (Ref : 15)	U07181 (SEQ ID NO's 32-33)	X51905	Y00711	Glycolysis	Oxidoreductas
Putative cytochrome c oxidase VIB (EC 1.9.3.1) (Ref : 16)			X13923 (SEQID NO's 34-35)	Mitochrondrial energy metabolism	Oxidoreductas
Putative NADH: ubiquinone oxidoreductasePG IV subunit (Ref: 17)			AF044953(SEQ ID No's 36-37)	Mitochrondrial energy metabolism	Oxidoreductas
Putative succinate dehydrogenase flavoprotein (Ref: N/A)		AF095938 (SEQ ID No's 38-39)	AF171022	TCA cycle	Oxidoreductas
Putative ubiquinol- cytochrome-c reductase (EC 1.10.2.2) core protein II (REF: 18)			J04973 (SEQ ID No's 40-41)	Mitochrondrial energy metabolism	Oxidoreductas
Stearoyl-coA desaturase 2 * (Ref: N/A)	AB032243 (SEQ ID N0's 42-43)	M26270		Fatty acid biosynthesis	Oxidoreductas
Ribophorin I * (Ref: 19)	X05300 (SEQ ID No's 44-45)	-		Glycosolation	Transferase
Sulfotransferase- like protein (REF : 20)	AF188699 (SEQ ID No's 46-47)			Transferase	Transferase
ATP synthase, H+, alpha subunit, mitochondrial (EC 3.6.1.34) (Ref: N/ A)	X56133 (SEQ ID No's 48-49)			Mitochrondrial energy metabolism	Hydrolase
F1F0 ATPase delta subunit (REF : 21)	U00926 (SEQ ID No's 50-51)			Mitochrondrial energy metabolism	Hydrolase
Putative dihydropyrimidin ase related protein * (REF: 22)		·	D78013 (SEQ ID NO's 52-53)	Pyrimidine degradation	Hydrolase
Heat shock protein 90 (REF : 23)	S45392 (SEQ ID NO's 114-115)	M18186	M16660	Cell protection	Hydrolase

Table VI - (continued)

Expression product or Name	Rat Accession Number	Mouse Accession Number	Human Accession Number	Reaction	Assay
Myelin basic protein S (MBP S) (REF :24)	K00512 SEQ ID No 54)			Myelin structural protein	Ligand binding
Transferrin (Ref 25)	D38380 (SEQ ID No's 55-56)			Iron transport	Ligand binding
Neurofilament, light molecular weight (NF-L) (Ref : 26)	AF031880 (SEQ ID No's 57-58)		·	Cytoskeleton	Ligand binding
Myelin- associayted glycoprotein (MAG) (Ref : 27)	M1680 (SEQ ID NO's 59-60)	M31811		Cell adhesion molecule for postnatal neural development	Ligand binding
NF-M middle molecular weight neurofilament protein (Ref : 28)	M18628 (SEQ ID No's 61-62)			Cytoskeleton	Ligand binding
Neuro- degeneration associated-protein 1 (Ref: 29)	D32249 (SEQ ID No's 63-64)			Protein sorting; synaptic communication and plasticity	Ligand binding
S-100 protein β- subunit (REF: 30)	X01090 (SEQ ID No 65)			Zinc and calcium binding	Ligand binding
Microtubule- associated protein 1b (Map 1b) (Ref: 31)	X60370 (SEQ ID No's 66-67)		L06237	Cytoskeleton protein, neuronal growth/ regeneration; microtubule binding protein	Ligand binding
Putative cdc 37 homolog (Ref: 32)	D26564 (Seq ID No's 68-69)			Cell signalling; cell cycle protein	Ligand binding
Putative ras- related protein Rab-5c (Ref : 33)			U11293 (SEQID No's 70-71)	Small GTP binding protein	Ligand binding
Putative gelsolin (Ref: 34)		J04953 (SEQ ID No's 72-73)		Cytoskeleton protein	Ligand binding
Cd81 antigen (target of antiproliferative antibody 1) (Ref : 35)	U19894 (SEQ ID No's 74-75)	X59047	M33680	Regulator for neuron-induced astrocyte differentiation; microglial effector functions	Ligand binding

Table VI - (continued)

ſ	Sequences whose e	expression produ	cts are other en	zymes		
5	Expression product or Name	Rat Accession Number	Mouse Accession Number	Human Accession Number	Reaction	Assay
10	Mobp81 (Myelin- associated/Oligo- dendrocytic basic protein 81) (Ref : 36)	X87900 (SEQ ID No's 76-77)			Myelin compaction	Ligand binding
15	Syntaxin binding protein n-sec1, sec1 homolog (Ref: N/A)			BC002869 (Seq ID No's 78-79)	Neuro- transmission	Ligand binding
20	Alpha-internexin (Ref: 37)	M73049 (SEQ ID No's 80-81)			Cytoskeleton protein; neuronal intermediate filament that can self-assemble	Ligand binding
25	Putative β- sarcoglycan A3b (Ref: N/a)		AB024921 (SEQ ID No's 82-83)		Cytoskeleton protein	Ligand binding
	CGI-78 protein Ref: 38)	AF151835		AF151835(SEQ ID No's 84-85)	Unknown	Ligand binding
<i>30</i>	KIAA0143 (Ref: 39)			D63477 (SEQ ID NO's 86-87)	Transmembrane protein	Ligand binding
	Septin 2 (KIAA0128 ; Ref 39)	,		D50918 (SEQ ID No's 88-89)	GTPase; cytokinesis	Ligand binding
35	Nucleobindin (Ref: 40)	Z36277 (SEQ ID No's 90-91)			Unknown	Ligand binding
	Myelin protein SR13 (Ref: 41)	M69139 (SEQ ID NO's 92-93)	S55427		Myelin structural protein	Ligand binding
40	B-Actin, cytoplasmic (Ref: 42)	V01217 (SEQ ID NO's 94-95)	X03672		Structural protein	Ligand binding
45	Ly6/neurotoxin (Lynx1) homolog (Ref: 43)		AF141377 (SEQ ID No's 96-97)		Neuro- transmitter modulator	Ligand binding
	Astrocytic phosphoprotein; PFA 15 gene (Ref: N/A)	AJ243949 (SEQ ID NO's 98-99)	X86694		PKC substrate	Ligand binding
50	PLIC-1 (Ref: N/A)		AF177345 (SEQ ID NO 100-101)		Cytoskeleton interaction	Ligand binding
55	Nfx1 (tip associating protein (TAP) gene) (Ref: N/A)	AF093139 (SEQ ID NO's 102-103)	AF093140		Unknown	Logand binding

Table VI - (continued)

Expression product or Name	Rat Accession Number	Mouse Accession Number	Human Accession Number	Reaction	Assay
Alpha-Crystallin B (Ref N/A)	U04320 SEQ ID NO's 104- 105)	M73741	M28638	Stress response; heat shock element	Ligand binding
Heat shock-like protein 70 kD (Ref : 44)	X70065 (SEQ ID No's 106-107)	U73744	Y00371	Stress response	Ligand binding
Tau microtubule- associated protein (Ref : 45)	X79321 (SEQ ID NO's 108-109)			Microtubule associated protein expressed in neurons	Ligand binding
Myelin, Schwann cell, Peripheral (P- 0) (Ref : 46)	K03242 (SEQ ID No's 110-111)			Myelin structural protein	Ligand binding
B-Tubulin class 1 (Ref: 47)	AB011679 (SEQ ID NO's 112-113)	X04663	AF14139	Structural protein	Ligand binding
Putative ribonuclease III (Ref : N/A)			AF116910 (SEQ ID NO's 116-117)	RNA hydrolysis	Hydrolase

PRODUCTION OF POLYPEPTIDES AND NUCLEIC ACIDS

Vectors

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[0017] Recombinant expression vectors comprising a nucleic acid can be employed to express any of the nucleic acid sequences of the invention. The expression products derived from such vector constructs can be used to develop screening technologies for the identification of molecules that can be used to prevent or treat pain, and in the development of diagnostic tools for the identification and characterization of pain. The expression vectors may also be used for constructing transgenic non-human animals.

[0018] Gene expression requires that appropriate signals be provided in the vectors, said signals including various regulatory elements such as enhancers/promoters from viral and/or mammalian sources that drive expression of the genes or nucleic acid sequences of interest in host cells. The regulatory sequences of the expression vectors used in the invention are operably linked to the nucleic acid sequence encoding the pain-associated protein of interest or a peptide fragment thereof.

[0019] Generally, recombinant expression vectors include origins of replication, selectable markers, and a promoter derived from a highly expressed gene to direct transcription of a downstream nucleoide sequence. The heterologous nucleotide sequence is assembled in an appropriate frame with the translation, initiation and termination sequences, and if applicable a leader sequence to direct the expression product into the periplasmic space, the extra-cellular medium or cell membrane.

[0020] In a specific embodiment wherein the vector is adapted for expressing desired sequences in mammalian host cells, preferred vectors will comprise an origin of replication from the desired host, a suitable promoter and an enhancer, and also any necessary ribosome binding sites, polyadenylation site, transcriptional termination sequences, and optionally 5'-flanking non-transcribed sequences. DNA sequences derived from the SV40 or CMV viral genomes, for example SV40 or CMV origin, early promoters, enhancers, and polyadenylation sites may be used to provide the required non-transcribed genetic elements.

[0021] A recombinant expression vector used in the invention advantageously also comprises an untranscribed poly-

nucleotide region located at the 3'end of the coding sequence (ORF), this 3'-UTR (untranslated region) polynucleotide being useful for stabilizing the corresponding mRNA or for increasing the expression rate of the vector insert if this 3'-UTR harbours regulation signal elements such as enhancer sequences.

[0022] Suitable promoter regions used in the expression vectors are chosen taking into account the host cell in which the nucleic acid sequence is to be expressed. A suitable promoter may be heterologous with respect to the nucleic acid sequence for which it controls the expression, or alternatively can be endogenous to the native polynucleotide containing the coding sequence to be expressed. Additionally, the promoter is generally heterologous with respect to the recombinant vector sequences within which the construct promoter/coding sequence has been inserted. Preferred promoters are the Lacl, LacZ, T3 or T7 bacteriophage RNA polymerase promoters, the lambda PR, PL and Trp promoters (see EP-0 036 776), the polyhedrin promoter, or the p10 protein promoter from baculovirus (kit Novagen; Smith et al., (1983); O'Reilly et al. (1992)).

[0023] Preferred selectable marker genes contained in the expression recombinant vectors used in the invention for selection of transformed host cells are preferably dehydrofolate reductase or neomycin resistance for eukaryotic cell culture, TRP1 for *S. cerevisiae* or tetracycline, rifampicin or ampicillin resistance in *E. coli*, or Levamsaccharase for *Mycobacteria*, this latter marker being a negative selection marker.

[0024] Preferred bacterial vectors are listed hereafter as illustrative but not limitative examples: pQE70, pQE60, pQE-9 (Quiagen), pD10, phagescript, psiX174, p.Bluescript SK, pNH8A, pNH16A, pNH18A, pNH46A (Stratagene); pKK223-3, pKK233-3, pDR540, pRIT5 (Pharmacia); pWLNEO, pSV2CAT, pOG44, pXT1, pSG (Stratagene); pSVK3, pBPV, pMSG, pSVL (Pharmacia); pQE-30 (QIA express).

[0025] Preferred bacteriophage recombinant vectors of the invention are P1 bacteriophage vectors such as described by Sternberg N.L. (1992;1994).

[0026] A suitable vector for the expression of any of the pain associated polypeptides used in the invention or fragments thereof, is a baculovirus vector that can be propagated in insect cells and in insect cell-lines. A specific suitable host vector system is the pVL 1392/1393 baculovirus transfer vector (Pharmingen) that is used to transfect the SF9 cell line (ATCC N°CRL 1711) that is derived from Spodoptera frugiperda.

[0027] The recombinant expression vectors of in the invention may also be derived from an adenovirus. Suitable adenoviruses are described by Feldman and Steig (1996) or Ohno *et al.* (1994). Another preferred recombinant adenovirus is the human adenovirus type two or five (Ad 2 or Ad 5) or an adenovirus of animal origin (Patent Application WO 94/26914)

[0028] Particularly preferred retrovirus for the preparation or construction of retroviral *in vitro* or *in vivo* gene delivery vehicles include retroviruses selected from the group consisting of Mink-Cell Focus Inducing Virus, Murine Sarcoma Virus, and Ross Sarcoma Virus. Other preferred retroviral vectors are those described in Roth *et al.* (1996), in PCT Application WO 93/25234, in PCT Application WO 94/06920, and also in Roux *et al.* (1989), Julan *et al.* (1992) and Nada *et al.* (1991).

[0029] Yet, another viral vector system that is contemplated is the Adeno Associated Viruses (AAV) such as those described by Flotte et al. (1992), Samulski et al. (1989) and McLaughlin et al. (1996).

Host cells expressing pain associated polypeptides

40 [0030] Host cells that endogenously express pain associated polypeptides or have been transformed or transfected with one of the nucleic acid ssequences described herein, or with one of the recombinant vector described above, particularly a recombinant expression vector, can be used in the present invention. Also included are host cells that are transformed (prokaryotic cells) or are transfected (eukaryotic cells) with a recombinant vector such as one of those described above.

45 [0031] Preferred host cells used as recipients for the expression vectors used in the invention are the following:

(a) prokaryotic host cells: Escherichia coli, strains. (i.e. DH5-α, strain) Bacillus subtilis, Salmonella typhimurium and strains from species like Pseudomonas, Streptomyces and Staphylococcus for the expression of up and down-regulated nucleic acid sequence modulated by pain, characterized by having at least 80% sequence identity with any of the nucleic acid sequences of Tables I - VI. Plasmid propagation in these host cells can provide plasmids for transfecting other cells.

(b) eukaryotic host cells: HeLa cells (ATCC N°CCL2; N°CCL2.1; N°CCL2.2), Cv 1 cells (ATCC N°CCL70), COS cells (ATCC N°CRL 1650; N°CRL 1651), Sf-9 cells (ATCC N°CRL 1711), C127 cells (ATCC N°CRL-1804), 3T3 cells (ATCC N°CRL-6361), CHO cells (ATCC N°CCL-61), human kidney 293 cells (ATCC N° 45504; N°CRL-1573), BHK (ECACC N°84100 501; N°84111301), PC12 (ATCC N° CRL-1721), NT2, SHSYSY (ATCC N° CRL-2266), NG108 (ECACC N°88112302) and F11, SK-N-SH (ATCC N° CRL-HTB-11), SK-N-BE(2) (ATCC N° CRL-2271), IMR-32 (ATCC N° CCL-127). A preferred system to which the nucleic acids of the invention can be expressed are

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neuronal cell lines such as PC12, NT2, SHSY5Y, NG108 and F11, SK-N-SH, SK-N-BE(2), IMR-32 cell lines, COS cells, 3T3 cells, HeLa cells, 292 cells and CHO cells. The above cell lines could be used for the expression of any of the nucleic acid sequences of Tables I - VI.

[0032] When a nucleic acid sequence of any of Tables I - X is expressed using a neuronal cell line, the sequence can be expressed through an endogenous promoter or native neuronal promoter, or an exogenous promoter. Suitable exogenous promoters include SV40 and CMV and eukaryotic promoters such as the tetracycline promoter. The preferred promoter when pain associated molecules are endogenously expressed is an endogenous promoter. A preferred promoter in a recombinant cell line is the CMV promoter.

[0033] In a specific embodiment of the host cells described above, these host cells have also been transfected or transformed with a polynucleotide or a recombinant vector for the expression of a natural ligand of any of the nucleic acid sequences of any of Tables I - VI or a modulator of these expression products.

PROTEINS, POLYPEPTIDES AND FRAGMENTS

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[0034] The expression products of the nucleic acid sequences of Tables I - VI or fragment(s) thereof can be prepared using recombinant technology, from cell lines or by chemical synthesis. Recombinant methods, chemical methods or chemical synthetic methods can be used to modify a gene in order to introduce into the gene product, or a fragment of the gene product, features such as recognition tags, cleavage sites or other modifications. For efficient polypeptide production, the endogenous expression system or recombinant expression system should allow the expression products to be expressed in a manner that will allow the production of a functional protein or fragment thereof which can be purified. Preferred cell lines are those that allow high levels of expression of polypeptide or fragments thereof. Such cell lines include cell lines which naturally express the nucleic acid sequence of Tables I - VI or common mammalian cell lines such as CHO cells or COS cells, etc, or more specific neuronal cell lines such as PC12. However, other cell types that are commonly used for recombinant protein production such as insect cells, amphibian cells such as oocytes, yeast and prokaryotic cell lines such as *E. coli* can also be used.

[0035] The expression products of Tables I - VI or fragments thereof can be utilized in screens to identify potential therapeutic ligands, either as a purified protein, as a protein chimera such as those produced in phage display, as a cell membrane (lipid or detergent) preparation, or in intact cells.

[0036] The invention also relates generally to the use of proteins, peptides and peptide fragments for the development of screening technologies for the identification of molecules for the prevention or treatment of pain, and the development of diagnostic tools for the identification and characterization of pain. These peptides include expression products of the nucleic acid sequences of Tables I - VI and purified or isolated polypeptides or fragments thereof having at least 90%, preferably 95%, more preferably 98% and most preferably 99% sequence identity with the any of the expression products of nucleic acid sequences of Tables I - VI. Expressed peptides and fragments of any of these nucleic acid sequences can be used to develop screening technologies for the identification of novel molecules for the prevention or treatment of pain. These screening technologies could also be used to ascribe new pain therapeutic indications to molecules, which have not previously been ascribed for the prevention or treatment of pain. Furthermore the said expressed peptides and fragments can be used as diagnostic tools and for the development of diagnostic tools.

SCREENING METHODS

[0037] As discussed above, we have identified nucleic acid sequences whose expression is regulated by pain, particularly chronic pain and more particularly diabetic pain. The expression products of these nucleic acids can be used for screening ligand molecules for their ability to prevent or treat pain, and particularly, but not exclusively, chronic pain. The main types of screens that can be used are described below. The test compound can be a peptide, protein or chemical entity, either alone or in combination(s), or in a mixture with any substance. The test compound may even represent a library of compounds.

[0038] The expression products of any of the nucleic acid sequences of Tables I - VI or fragments thereof can be utilized in a ligand binding screen format, a functional screen format or invivo format. Examples of screening formats are provided.

A) Ligand binding screen

[0039] In ligand binding screening a test compound with is contacted with an expression product of one of the sequences of Tables I - VI, and the ability of said test compound to interact with said expression product is determined, e.g. the ability of the test compound(s) to bind to the expression product is determined. The expression product can be a part of an intact cell, lipidic preparation or a purified polypeptide(s), optionally attached to a support, such as

beads, a column or a plate etc.

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[0040] Binding of the test compound is preferably performed in the presence of a ligand to allow an assessment of the binding activity of each test compound. The ligand may be contacted with the expression product either before, simultaneously or after the test compound. The ligand should be detectable and/or quantifiable. To achieve this, the ligand can be labelled in a number of ways, for example with a chromophore, radioactive, fluorescent, phosphorescent, enzymatic or antibody label. Methods of labelling are known to those in the art. If the ligand is not directly detectable it should be amenable to detection and quantification by secondary detection, which may employ the above technologies. Alternatively the expression product or fragment thereof can be detectable or quantifiable. This can be achieved in a similar manner to that described above.

[0041] Binding of the test compound modifies the interaction of the ligand with its binding site and changes the affinity or binding of the ligand for/to its binding site. The difference between the observed amount of ligand bound relative to the theoretical maximum amount of ligand bound (or to the ligand bound in the absence of a test compound under the same conditions) is a reflection of the binding ability (and optionally the amount and/or affinity) of a test compound to bind the expression product.

[0042] Alternatively, the amount of test compound bound to the expression product can be determined by a combination of chromatography and spectroscopy. This can be achieved with technologies such as Biacore (Amersham Pharmacia). The amount of test compound bound to the expression product can also be determined by direct measurement of the change in mass upon compound or ligand binding to the expression product. Alternatively, the expression product, compound or ligand can be fluorescently labelled and the association of expression product with the test compound can be followed by changes in Fluorescence Energy Transfer (FRET).

[0043] The invention therefore includes a method of screening for pain alleviating compounds, comprising:

- a) contacting a test compound or test compounds in the presence of a ligand with an expression product of any of the nucleic acid sequences of Tables I VI or with a cell expressing at least one copy of the expression product or with a lipidic matrix containing at least one copy of the expression product:
- b) determining the binding of the test compound to the expression product, and
- c) selecting test compounds on the basis of their binding abilities.

[0044] In the above method, the ligand may be added prior to, simultaneously with or after contact of the test compound with the expression product. Non limiting examples and methodology can be gained from the teachings of the Molecular Probes handbook and references therein (Molecular Probes, Inc., 4849 Pitchford Ave, Eugene, USA), Methods in neurotransmitter receptor analysis (Yamamura HI., Enna, SJ., and Kuhar, MJ., Raven Press New York, the Glaxo Pocket Guide to Pharmacology, Dr. Michael Sheehan, Glaxo Group Research Ltd, Ware, Herts SG12 0DP), Bylund DB and Murrin LC (2000, Life Sciences, 67 (24) 2897-911), Owicki JC (2000, J. biomol Screen (5) 297-306), Alberts et al (1994, Molecular Biology of the Cell, 3rd Edn, Garland Publication Inc), Butler JE, (2000 Methods 22(1):4-23, Sanberg SA (2000, Curr Opin Biotechnol 11(1) 47-53), and Christopoulos A (1999, biochem Pharmacol 58(5) 735-48).

B) Functional screening

(a) Kinase assays

[0045] The expression product of any of the nucleic acid sequences of Tables I - VI which encode a kinase, and in particular the nucleic acid sequence listed in Table I, is amenable to screening using kinase assay technology.

[0046] Kinases have the ability to add phosphate molecules to specific residues in ligands such as binding peptides in the presence of a substrate such as adenosine triphosphate (ATP). Formation of a complex between the kinase, the ligand and substrate results in the transfer of a phosphate group from the substrate to the ligand. Compounds that modulate the activity of the kinase can be determined with a kinase functional screen. Functional screening for modulators of kinase activity therefore involves contacting one or more a test compounds with an expression product of one of the nucleic acid sequences of Tables I - VI which encodes a kinase, and determining the ability of said test compound to modulate the transfer of a phosphate group from the substrate to the ligand.

[0047] The expression product can be part of an intact cell or of a lipidic preparation or it can be a purified polypeptide (s), optionally attached to a support, for example beads, a column, or a plate. Binding is preferably performed in the presence of ligand and substrate to allow an assessment of the binding activity of each test compound.

[0048] The ligand should contain a specific kinase recognition sequence and it should not be phosphorylated at its phosphoryation site. The ligand and/or substrate may be contacted with the kinase either before, simultaneously or after the test compound. Optionally the substrate may be labelled with a kinase transferable labelled phosphate. The assay being monitored by the phosphorylation state of the substrate and/or the ligand. The ligand should be such that its phosphorylation state can be determined. An alternative method to do this is to label the ligand with a phosphorylation.

ylation-state-sensitive molecule. To achieve this, the ligand can be labelled in a number of ways, for example with a chromophore, radioactive, fluorescent, phosphorescent, enzymatic or antibody label. If the ligand is not directly detectable it should be amenable to detection and quantification by secondary detection, which may employ the above technologies. Such technologies are known to those in the art.

[0049] Binding of the test compound to the kinase modifies its ability to transfer a phosphate group from the substrate to the ligand. The difference between the observed amount of phosphate transfer relative to the theoretical maximum amount of phosphate transfer is a reflection of the modulatory effect of the test compound. Alternatively, the degree of phosphate transfer can be determined by a combination of chromatography and spectroscopy. The extent of phosphorylation of the ligand peptide or dephosphorylation of the substrate can also be determined by direct measurement. This can be achieved with technologies such as Biacore (Amersham Pharmacia).

[0050] The invention also provides a method for screening compounds for the ability to relieve pain, which comprises:

- (a) contacting one or more test compounds in the presence of ligand and substrate with an expression product of any of the sequences of Tables I VI which is a kinase or with a cell containing at least 1 copy of the expression product or with a lipidic matrix containing at least 1 copy of an expression product;
- (b) determining the amount of phosphate transfer from the substrate to the ligand; and
- (c) selecting test compounds on the basis of their capacity to modulate phosphate transfer.

[0051] Optionally ligand, substrate and/or other essential molecules may be added prior to contacting the test compound with expression product of step (a) or after step (a). Non limiting examples and methodology can be gained from the teachings of the *Molecular Probes handbook* and references therein (Molecular Probes, Inc., 4849 Pitchford Ave, Eugene, USA), *Methods in Molecular Biology 2000*; 99: 191-201, *Oncogene* 2000 20; 19(49): 5690-701, and *FASAB Journal*, (10, 6, P55, P1458, 1996, Pocius D Amrein K *et al*).

b) Receptor assays

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[0052] An expression product of any of the nucleic acid sequences of Tables I - VI which encodes a receptor, and in particular any of the nucleic acid sequences listed in Table II, is amenable to screening using receptor assay technology.

[0053] Receptors are membrane associated proteins that initiate intracellular signalling upon ligand binding. Therefore, the identification of molecules for the prevention and treatment of pain can be achieved with the use of a ligand binding assay, as outlined above. Such an assay would utilize an endogenous or non-endogenous ligand as a component of the ligand binding assay. The binding of this ligand to the receptor in the presence of one or more test compounds would be measured as described above. Such is the nature of receptors that the assay is usually, but not exclusively performed with a receptor as an intact cell or membraneous preparation.

[0054] The invention therefore includes a method of screening for pain alleviating compounds, comprising:

- a) contacting a test compound or test compounds in the presence of a ligand with a cell expressing at least one copy of the expression product of any of the sequences of Tables I VI which is a receptor or with a lipidic matrix containing at least one copy of the expression product;
- b) determining the binding of the test compound to the expression product, and
- c) selecting test compounds on the basis of their binding abilities.

c) Transporter protein assays

[0055] An expression product of any of the nucleic acid sequences of Tables 1 - VI which encodes a transporter protein, and in particular any of the nucleic acid sequences listed in Table III, is amenable to screening using transporter protein assay technology. Non limiting examples of technologies and methodologies are given by Carroll FI, et al (1995, Medical Research Review, Sep15 (5) p419-444), Veldhuis JD and Johnson MI (1994, Neurosci. Biobehav Rep., winter 18(4) 605-12), Hediger MA and Nussberger S (1995, Expt Nephrol, July-Aug 3(4) p211-218, Endou H and Kanai Y, (1999, Nippon Yakurigaku Zasshi, Oct. 114 Suppl 1:1p-16p), Olivier B et al (2000, Prog. Drug Res., 54, 59-119), Braun A et al (2000, Eur J Pharm Sci, oct 11, Suppl 2 S51-60) and Molecular Probes handbook and references therein (Molecular Probes, Inc., 4849 Pitchford Ave, Eugene, USA).

[0056] The main function of transporter proteins is to facilitate the movement of molecules across a cellular membrane. Compounds that modulate the activity of transporter proteins can be determined with a transporter protein functional screen. Functional screening for modulators of transporter proteins comprises contacting at least one test compound with an expression product as aforesaid which is a transporter protein and determining the ability of said test compound to modulate the activity of said transporter protein. The expression product can be part of an intact cell,

or lipidic preparation, optionally attached to a support, for example beads, a column or a plate. Binding is preferably performed in the presence of the molecule to be transported, which should only able to pass through a cell membrane or lipidic matrix with the aid of the transporter protein. The molecule to be transported should be able to be followed when it moves into a cell or through a lipidic matrix. Preferably the molecule to be transported is labelled to aid in characterization, e.g. with a chromophore, radioactive, fluorescent, phosphorescent, enzymatic or antibody label. If the molecule to be transported is not directly detectable it should be amenable to detection and quantification by secondary detection, which may employ the above technologies. The molecule to be transported may be contacted with the transporter protein before, simultaneously with or after the test compound. If binding of the test compound to the transporter protein modifies its ability to transport molecules through a membraneous or lipidic matrix, then the difference between the observed amount of transported molecule in a cell/or through a lipidic matrix relative to the theoretical maximum amount is a reflection of the modulatory effect of the test compound.

[0057] The invention further provides a method for screening compounds for their ability to relieve pain, comprising

- a) contacting at least one test compound in the presence of transporter molecules with a cell containing at least one copy of an expression product of any of the sequences of Tables I VI which is a transporter protein or with a lipidic matrix containing at least one copy of the expression product;
- c) measuring the movement of transported molecules into or from the cell, or across the lipidic matrix; and
- d) selecting test compounds on the basis of their ability to modulate the movement of transported molecules.

d) G-protein coupled receptor protein assays

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[0058] An expression product of any of the nucleic acid sequences of Tables I - VI that encodes a G-protein coupled receptor protein, and in particular any of the nucleic acid sequences listed in Table IV, is amenable to screening using G-protein coupled receptor protein assay technology.

[0059] G-protein coupled receptor proteins (GPCRs) are membrane associated proteins whose main function is to transduce a signal through a cellular membrane. Upon ligand binding, GPCRs undergo a conformational change that allows complexing of the GPCRs with a G-protein. G-proteins possess a GTP/GDP binding site. The formation of the G-protein/ligand complex allows exchange of GTP for GDP, resulting in a conformational change of the G-protein. This conformational change initiates signal transduction.

[0060] Functional screening for modulators of GPCRs comprises contacting at least one test compound with an expression product as aforesaid which is a G-protein coupled receptor protein, and assessing the ability of the test compound(s) to modulate the exchange of GTP for GDP or the modulation of the GPCR signal transduction pathway. The expression product can be part of an intact cell or lipidic preparation, optionally attached to a support, for example beads, a column or a plate. Binding is preferably performed in the presence of a ligand, G-protein and GTP/GDP to allow an assessment of the binding activity of each test compound. Alternatively components of the signaling pathway are also included. In particular, in a preferred embodiment, a labeled GTP is used and the ability of the test compound (s) to modulate the exchange of GTP to GDP is determined. A further optional characteristic of the assay can be the inclusion of a reporter molecule that enables monitoring the regulation of the signaling pathway. The ligand may be contacted with the GPCR before, simultaneously with, or after the test compound. Binding of the test compound to the GPCR modifies its ability to modulate the exchange of GTP for GDP and hence the modulation of signal transduction. The difference between the observed amount of GTP exchanged for GDP relative to the theoretical maximum amount of GTP is a reflection of the modulatory effect of the test compound. Likewise the relative activities of signal transduction reporter molecules are also a reflection of the modulatory effect of the test compound. Non limiting examples of technologies and methodologies can be found in Molecular Probes handbook and references therein (Molecular Probes, Inc., 4849 Pitchford Ave, Eugene, USA), Glaxo Pocket Guide to Pharmacology, (Michael Sheehan, Pharmacology Division staff, Glaxo Group Research Ltd., Ware, Herts SC12 0DP) and Xing et al (2000, J. Recept. Signal. Transduct. Res. 20(40 189-210).

[0061] The invention provides a method of screening compounds for their ability to relieve pain, comprising:

- a) contacting at least one test compound in the presence of a ligand, GTP/GDP, G-protein with a cell containing at least one copy of an expression product of a sequence of Tables I VI as aforesaid which is a G-protein coupled receptor protein or with a lipidic matrix containing at least one copy of the expression product;
- b) measuring the exchange of GTP for GDP, and
- c) selecting test compounds on the basis of their ability to modulate said exchange.

In the above method, the ligand, GTP/GDP, G-protein and other essential molecules can be added before, simultaneously with or after the contacting of the test compound(s) with the cell line or lipidic matrix in step (a).

e) DNA-binding protein assays

[0062] An expression product of any of the nucleic acid sequences of Tables I - VI that encodes a DNA-binding protein, and in particular any of the nucleic acid sequences listed in Table V, is amenable to screening using DNA-binding protein assay technology.

[0063] DNA binding proteins are proteins that are able to complex with DNA. The complexing of the DNA binding protein with the DNA in some instances requires a specific nucleic acid sequence. Screens can be developed in a similar manner to ligand binding screens as previously indicated and will utilise DNA as the ligand. DNA-binding protein assays can be carried using similar principles described in ligand binding assays as described above. Non limiting examples of methodology and technology can be found in the teachings of *Haukanes BI and Kvam* C (Biotechnology, 1993 Jan 11 60-63), *Alberts B et al* (Molecular Biology of the Cell, 1994, 3rd Edn., Garland Publications Inc, *Kirigiti P and Machida CA* (2000 Methods Mol Biol, 126, 431-51) and *Molecular Probes handbook* and references therein (Molecular Probes, Inc., 4849 Pitchford Ave., Eugene, USA).

[0064] The invention therefore includes a method of screening for pain alleviating compounds, comprising:

a) contacting a test compound or test compounds in the presence of a plurality of nucleic acid sequences with an expression product of any of the nucleic acid sequences of Tables I - VI which is a DNA binding protein or with cells expressing at least one copy of the expression product or with a lipidic matrix containing at least one copy of the expression product;

- b) determining the binding of the test compound to the expression product, and
- c) selecting test compounds on the basis of their binding abilities.
- In the above method, the plurality of nucleic acid sequence may be added prior to, simultaneously with or after contact of the test compound with the expression product.

f) Assays using other enzymes

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[0065] Expression products of any of the nucleic acid sequences of Tables I - VI that encode other enzymes, e.g. ligases, lyases, oxidoreductases, transferases and hydrolases, and in particular any of the nucleic acid sequences listed in Table VI, is amenable to screening using appropriate assay technology.

[0066] Each class of enzyme has a defined function. Ligases have the property of being able to splice molecules together. This is achieved with the conversion of ATP substrate to AMP. Therefore, the activity of a ligase can be followed by monitoring the conversion of ATP to AMP. Such technologies are known to those in the art. Non limiting examples and methodologies are illustrated by *Ghee. T. Tan et al* (1996,Biochem J. 314, 993-1000, *Yang SW et al* (1992, 15: 89(6) 2227-31 and refrences therein, and in *Molecular Probes handbook* and references therein (Molecular Probes, Inc., 4849 Pitchford Ave, Eugene, USA).

[0067] The invention also provides a methods of screening for pain alleviating compounds, comprising;

- a) contacting one or more test compounds in the presence of ATP with an expression product of any of the sequences of Tables I VI which is a ligase or with a cell expressing at least 1 copy of a expression product or with a lipidic matrix containing at least 1 copy of an expression product which is a ligase;
- b):determining the amount of ATP converted to AMP, and
- c) selecting test compounds on the basis of their ability to modulate said conversion.

[0068]. Lyases are enzymes which catalyse the cleavage of by reactions other than hydrolysis. These enzymes can be grouped into seven groups according to type of bond cleaved. These groups are carbon-carbon lyases (E.C. No 4.1), carbon-oxygene lyases (E.C. No 4.2), carbon-nitrogen lyases (E.C. No 4.3), carbon-sulphur lyase (E.C. No 4.4) carbon-halide lyases (E.C. No 4.5), phosphorous-oxygene lyases (E.C. No 4.6) and other lyases (E.C. No 4.99) (Analytical Biochemistry 3rd Edn, David J. Holme and Hazel Peck, Longman press). The enzyme commission number (E.C.) of the International Union of Biochemistry relates to the type of reaction catalysed by the enzyme. Further teachings on how to develop assays and screens for lyases can be obtained from Methods in Enzymology (Academic Press).

[0069] Oxidoreductases are enzymes that catalyse the transfer of hydrogen or oxygen atoms or electrons. These enzymes can by sub-grouped into twenty categories according to their specific mode of action. These groups are oxidoreductases acting on the CH-OH group of donors (E.C. No1.1 oxidoreductases acting on the aldehyde or oxo group of donors (E.C. No 1.2), oxidoreductases acting on the CH-NH2 group of donors (E.C. 1.4), oxidoreductases acting on the CH-NH group of donor (E.C. 1.5), oxidoreductases acting on the NADH or NADPH (E.C. No 1.6), oxidoreductases acting on other nitrogen compounds as donors (E.C. No 1.7), oxidoreductases acting on a sulphur group of donors (E.C. No 1.8), oxidoreductases acting on a haem group of donors (E.C. No 1.9), oxidoreductases acting on diphenols and related substances as donors (E.C. No 1.9), oxidoreductases acting on diphenols and related substances as donors (E.C.

C. 1.10), oxidoreductases acting on hydrogen peroxide as acceptor (E.C. No 1.11), oxidoreductases acting on hydrogen as donor (E.C. No 1.12), oxidoreductases acting on single doners with incorporation of molecular oxygen (E.C. No 1.13), oxidoreductases acting on paired donors with incorporation of molecular oxygen (E.C. No 1.14), oxidoreductases acting on superoxide radicals as acceptors (E.C. 1.15), oxidoreductases oxidizing metal ions (E.C. No 1.16), oxidoreductases acting on -CH2 groups (E.C. No 1.17), oxidoreductases acting on reduced ferredoxin as donor (E.C. No 1.18), oxidoreductases acting on reduced flavodoxin as donor (E.C. No 1.19) and other oxidoreductases (E.C. No 1.97) (Analytical Biochemistry 3rd Edn, David J. Holme and Hazel Peck, Longman Press). The enzyme commission number (E.C.) of the International Union of Biochemistry relates to the type of reaction catalysed by the enzyme. Further teachings on how to develop assays and screens for oxidoreductases can be obtained from *Methods in Enzymology* (Academic Press) with special reference to volume 249.

[0070] Transferases are enzymes that catalyse the transfer of specific groups. They are classified into eight sub groups according to function, transferring one-carbon group (E.C. No 2.1), Transfering aldehyde or ketonic residues (E.C. No 2.2), Acetyltransferases (E.C. 2.3), glycosyltransferases (E.C. No 2.4), transferring alkyl or aryl groups other than methyl groups (E.C. No 2.5), transferring nitrogeneous groups (E.C. No 2.6), transferring phosphorous-containing groups (E.C. No 2.7) and transferring sulphur-containing groups (E.C. No 2.8) (Analytical Biochemistry 3rd Edn, David J. Holme and Hazel Peck, Longman Press). The enzyme commission number (E.C.) of the International Union of Biochemistry relates to the type of reaction catalysed by the enzyme. Further teachings on how to develop assays and screens for transferases can be obtained from Methods in Enzymology (Academic Press).

[0071] Hydrolases are enzymes that catalyse hydrolytic reactions and are sub-grouped into eleven classes according to the type of reaction they carry out. Hydrolases acting on ester bonds (E.C. No 3.1), hydrolases acting on glycosyl compounds (E.C. No 3.2), hydrolases acting on ether bonds (E.C. No 3.3), hydrolases acting on peptide bonds (E.C. No 3.4), hydrolases acting on carbon-nitrogen bonds, other than peptide bonds (E.C. No 3.5), hydrolases acting on acid anhydrides (E.C. No 3.6), hydrolases acting on carbon-carbon bonds (E.C. No 3.7), hydrolases acting on halide bonds (E.C. No 3.8), hydrolases acting on phosphorous-nitrogen bonds (E.C. No 3.9), hydrolases acting on sulphur-nitrogen bonds (E.C. No 3.10) and hydrolases acting on carbon-phosphorous bonds (Analytical Biochemistry 3rd Edn, David J. Holme and Hazel Peck, Longman Press). The enzyme commission number (E.C.) of the International Union of Biochemistry relates to the type of reaction catalysed by the enzyme. Further teachings on how to develop assays and screens for hydrolases can be obtained from Methods in Enzymology (Academic Press) with special reference to volume 249.

C) In vivo functional screen

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[0072] Any of the nucleotide sequences described in Tables I - VI or homologues thereof may be inserted by means of an appropriate vector into the genome of a lower vertebrate or of an invertebrate animal or may be inactivated or down regulated in the genome of said animal. The resulting genetically modified animal may be used for screening compounds for effectiveness in the regulation of pain. The invertebrate may, for example, be a nematode e.g. Caenorhabditis elegans, which is a favourable organism for the study of response to noxious stimuli. Its genome sequence has been determined, see Science, 282, 2012 (1998), it can be bred and handled with the speed of a microorganism (it is a self-fertilizing hermaphrodite) and can therefore be used in a high throughput screening format (WO 00/63424, WO 00/63425, WO 00/63426 and WO 00/63427), and it offers a full set of organ systems, including a simple nervous system and contains many similarly functioning genes and signaling pathways to mammals. A thermal avoidance model based on a reflexive withdrawal reaction to an acute heat stimulus has been described by Wittenburg et al, Proc. Natl. Acad. Sci.USA, 96, 10477-10482 (1999), and allows the screening of compounds for the treatment of pain with the modulation of pain sensation as an endpoint.

[0073] The genome of *C. elegans* can be manipulated using homologous recombination technology which allows direct replacement of nucleic acids encoding *C. elegans* with their identified mammalian counterpart. Replacement of these nucleic acids with those nucleic acids outlined above would allow for the direct screening of test compound(s) with their expression products. Any of the pain-related genes described above may be ligated into a plasmid and introduced into oocytes of the worm by microinjection to produce germline transformants. Successful plasmid injection into *C. elegans* and expression of inserted sequences has been reported by Devgen B.V., Ghent, Belgium. It is also possible to produce by routine methods worms in which the target sequences are down-regulated or not expressed (knock-out worms). Further non limiting examples of methodology and technology can be found in the teachings of *Hazendonk et al* (1997, Nat genet. 17(1) 119-21), *Alberts et al*, (1994, Molecular Biology of the Cell 3rd Ed. Garland Publishing Inc, *Caenorhabditis elegans* is anatomically and genetically simple), *Broverman S et al*, (1993, PNAS USA 15;90(10) 4359-63) and Mello et al (1991, 10(12)3959-70).

[0074] A further method for screening compounds for ability to modify response to pain, e.g. relieve pain, comprises:

(a) contacting one or more test compounds with at least one C elegans containing at least one copy of a sequence

as set out above;

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- (b) subjecting the C. elegans to a nociceptive stimulus;
- (c) observing the response of the C. elegans to said stimulus; and
- d) selecting test compounds on the basis of their ability to modify the response of C. elegans to said stimulus.

DIAGNOSTIC TOOLS AND KITS

Affinity peptides, ligands and substrates

[0075] Pain associated polypeptides and fragments thereof can be detected at the tissue and cellular levels with the use of affinity peptides, ligands and substrates, which will enable a skilled person to define more precisely a patient's ailment and help in the prescription of a medicament. Such affinity peptides are characterized in that firstly they are able to bind specifically to a pain associated polypeptide, and secondly that they are capable of being detected. Such peptides can take the form of a peptide or polypeptide for example an antibody domain or fragment, or a peptide/ polypeptide ligand or substrate, or a polypeptide complex such as an antibody.

[0076] The preparation of such peptides and polypeptides are known to those in the art. Antibodies, these may be polyclonal or monoclonal, and include antibodies derived from immunized animals or from hybridomas, or derivatives thereof such as humanized antibodies, Fab or F(ab')2 antibody fragments or any other antibody fragment retaining the antigen binding specificity.

[0077] Antibodies directed against pain-associated gene product molecules may be produced according to conventional techniques, including the immunization of a suitable mammal with the peptides or polypeptides or fragment thereof. Polyclonal antibodies can be obtained directly from the serum of immunized animals. Monoclonal antibodies are usually produced from hybridomas, resulting from a fusion between splenocytes of immunized animals and an immortalized cell line (such as a myeloma). Fragments of said antibodies can be produced by protease cleavage, according to known techniques. Single chain antibodies can be prepared according to the techniques described in US 4,946,778. Detection of these affinity peptides could be achieved by labeling, technologies which allow detection of peptides, such as enzymatic labeling, fluorescence labeling or radio-labeling are well known to those in the art. Optionally these affinity peptides, ligands and substrates could themselves be detected with the use of a molecule that has specific affinity to the peptides, ligands and substrates and is itself labeled.

30 [0078] The invention further provides a kit comprising;

- (a) affinity peptide and/or ligand and/or substrate for an expression product of a gene sequence that is down-regulated in the spinal cord of a mammal in response to first and second models of neuropathic or central sensitization pain; and
- (b) a defined quantity of an expression product of a gene sequence that is down-regulated in the spinal cord of a mammal both in response to first and second models of neuropathic or central sensitization pain,

for simultaneous, separate or sequential use in detecting and/or quantifying an expression product of a gene sequence that is down-regulated in the spinal cord of a mammal in response to first and second models of neuropathic or central sensitization pain.

Complimentary nucleic acids

[0079] Pain associated nucleic acid sequences can be characterized at the tissue and cellular levels with the use of complimentary nucleic acid sequences. Detection of the level of expression of pain associated nucleic acid sequences can help in the prognosis of a pain condition and the prescription of a medicament. These complimentary nucleic acids are characterized in that they can hybridize to a pain associated nucleic acid sequence and their presence can be detected through various techniques. Such techniques are known to those in the art and may include detection by polymerase chain reaction or detection by labeling of complimentary nucleic acid sequences by enzymatic labeling, affinity labeling fluorescent labeling or radio labeling. Complimentary strand nucleic acid sequences of this invention are 10 to 50 bases long, more preferably 15 to 50 bases long and most preferably 15 to 30 bases long, and hybridize to the coding sequence of the nucleic acid sequence.

[0080] A further aspect of this invention is a kit that comprises:

(a) nucleic acid sequences capable of hybridization to a nucleic acid sequence that is down-regulated in the spinal cord of a mammal in response to first and second models of neuropathic or central sensitization pain; and
 (b) a defined quantity of one or more nucleic acid sequences capable of hybridization to a nucleic acid sequence that is down-regulated in the spinal cord of a mammal in response to first and second models of neuropathic or

central sensitization pain, for simultaneous, separate or sequential use in detecting and/or quantifying a gene sequence that is down-regulated in the spinal cord of a mammal in response to first and second models of neuropathic or central sensitization pain.

5 IDENTIFICATION AND VALIDATION

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[0081] Subtractive hybridization enables the identification of nucleic acid sequences whose expression profiles are modified by a stimulus. Upon stimulation of a system (in the case of this invention a nociceptive stimulus on an animal model) all observed changes in the level of nucleic acid sequence expression are due to the reaction of the system to the stimulus. Characterization of these changes in expression by way of identification of nucleic acid sequence and level of expression is both identification and validation.

[0082] The inventors have developed a four step process which allows for the simultaneous identification and validation of nucleic acid sequences whose expression are regulated by a pain stimulus, preferably a chronic pain stimulus, and more preferably a diabetic pain stimulus. This process may comprise the following steps:

(a) induction of a nociceptive stimulus in test animals;

- (b) extraction of nucleic acids from specific neuronal tissue of test and control animals;
- (c) selective amplification of differentially expressed nucleic acid sequences; and
- (d) identification and characterization of differentially expressed gene products that are modulated by a nociceptive stimulus.

[0083] The above process is described in more detail below.

(a) Induction of nociceptive stimulus

[0084] The effect of the selected nociceptive stimulus on the test animal needs to be confirmable. The test subjects are therefore a species that has a "developed" nervous system, preferably similar to that of humans, most preferably rats or mice. Advantageously, the nociceptive stimulus is analogous to known pain paradigms in humans. One such paradigm of pain is the pain associated with diabetes, which can be induced in rodents with the use of streptozotocin (STZ). The present application requires the sequences to be down-regulated in two pain models which may be, but are not limited to models of neuropathic pain and/or central sensitization, and in which diabetic pain provides the first model and mechanical damage e.g. to a nerve leading into the spine can provide an appropriate second model.

[0085] Streptozotocin (STZ) induces hyperglycemia and Type 1 diabetes mellitus in rats. In particular, STZ contains a glucose analogue that allows it to be taken up by the glucose transporter 2 present on the surface of pancreatic β cells, the site of insulin synthesis. Once inside the cell, STZ causes a reduction in the level of nicotinamide adenine dinucleotide (NAD+). The decrease in NAD+ levels eventually leads to necrosis of the pancreatic β cell, causing a reduction in insulin levels and then diabetes, leading to neuropathy (diabetic) and neuropathic pain (R. B. Weiss, *Cancer Treat. Rep.*, 66, 427-438 1982, Guy *et al*, *Diabetologica*, 28, 131-137 1985; Ziegler *et al*, *Pain*, 34, 1-10 1988; Archer *et al*, *J. Neural. Neurosurgeon. Psychiatry*, 46, 491-499 1983). The diabetic rat model has been shown to be a reliable model of hyperalgesia. We have used an STZ-induced diabetic rat model to create a state of hyperlagesia that can be compared with control animals (Courteix *et al*, *Pain*, 53, 81-88 1993).

[0086] Three models of neuropathic and/or central sensitization pain in rats, which involve nerve injury, may be used, see Ralston, DD (1998) Present models of neuropathic pain. Pain Rev. 5: 83-100. The injuries are caused by (1) loosely tying four chromic gut sutures around the sciatic nerve (CCI model developed by Bennett, GJ and Y-K Xie, A peripheral mononeuropathy in rat that produces disorders of pain sensation like those seen in man, Pain 33: 87-107 (1988), (2) tightly ligating one third to one half of the fibers in the sciatic nerve (model developed by Seltzer, Z, R Dubner, Y Shire, A novel behavioral model of neuropathic pain disorders produced in rats by partial sciatic nerve injury, Pain 43: 205-218, 1990), and (3) tightly ligating the dorsal spinal nerve of a rat at the L5 or L5 and L6 levels (L5 model developed by Kim, SH and JM Chung, An experimental model for peripheral neuropathy produced by segmental spinal nerve ligation in the rat, Pain 50:355-363, 1992).

(b) Extraction of nucleic acids from neuronal tissue of test and control animals

[0087] The inventors have determined that RNA extraction of whole spinal cord nervous tissue would provide a way of identifying nucleic acid sequences whose expression in spinal tissue is modulated by streptozotocin induced diabetes or by a mechanical nerve damage model for neuropathic and/or central sensitization pain e.g. CCI. Test (subjected to the nociceptive stimulus) and control animals were sacrificed, and the tissue to be studied e.g. neural tissue separated. Techniques for so doing vary widely from animal to animal and will be familiar to skilled persons.

[0088] A cDNA library can be prepared from total RNA extracted from neural tissue of the test and control animals. Where possible, however, it is preferred to isolate the mRNA from the total RNA of the test and control animals, by affinity chromatography on oligo (dT)-cellulose, and then reverse transcribe the mRNA from the test and control animals to give test and control cDNA. Converting mRNA from the test and control animals to corresponding cDNA may be carried out by any suitable reverse transcription method, e.g. a method as described by Gubler & Hoffman, Gene, 25, 263-269 (1983). If desired a proprietary kit may be used e.g. the CapFinder PCR cDNA Library Construction Kit (Life Technologies) which is based on long-distance PCR and permits the construction of cDNA libraries from nanograms of total RNA.

(c) Selective amplification of differentially expressed nucleic acids

[0089] The reverse transcribed cDNA of the test and control animals is subjected to subtractive hybridisation and amplification so that differentially expressed sequences become selectively amplified and commonly expressed sequences become suppressed, so as to over-produce DNA associated with said nociceptic stimulus. A wide range of subtractive hybridisation methods can be used, but the preferred method is so-called suppression subtractive hybridisation, see US-A-5565340 and Diatchenko *et al, Proc. Nat. Acad. Sci. USA*, 93, 6025-6030 (1996), the disclosures of which are herein incorporated by reference. Kits for carrying out this method are available from CLONTECH Laboratories, Inc.

(d) Cloning and Sequencing the differentially expressed cDNA

[0090] The differentially expressed cDNA is ligated into a cloning vector, after which cells of *E. coli* are transformed with the vector and cultured. Positive clones are selected and lysed to release plasmids containing the cDNA insert. The plasmids are primed using forward and reverse primers to either side of the cloning site and the cDNA insert is sequenced. Vector and adaptor sequences are then removed from the output data from the sequencer, leaving only the nucleotide sequence of the differentially expressed gene. The sequence is then checked against data held in a database for homology to known nucleotide sequences and genes, including expressed sequence tags (ESTs) and coding sequences for proteins.

(e) Validation of the above method

[0091] The importance of the sequences that we have identified in pain is confirmed by the fact that genes have been identified using this method that represent nucleic acid sequences which have previously been implicated in pain, including Calmodulin (pRCM1, Genebank X13933), Enkephalin (Genebank Y07503) and Neurotensin receptor type 2 (Genebank X97121).

[0092] The inventors have identified nucleic acid sequences of the MAP kinase pathway, a previously non pain-associated biological pathway. The inventors have subsequently shown that intra-spinal injection of a MEK inhibitor (MEK is part of the MAP kinase pathway) produces a powerful inhibition of pain (Patent application No US 60/144292). Subsequently, it has been shown that the MAP kinase is also implicated in acute inflammatory pain (Woolf *et al*, Nature Neuroscience 1999).

[0093] The invention will now be further described in the following Example.

EXAMPLE

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45 Induction of diabetes

[0094] Diabetes was induced in adult (150-200g) male Sprague-Dawley rats (n=6) as described by Courteix *et al* (supra). Animals were injected intraperitoneally with streptozotocin (STZ)(50 mg/kg) dissolved in distilled water. Control or sham animals (age-matched animals, n=6) were injected with distilled water only.

Chronic Constrictive Injury (CCI)

[0095] Rats were anaesthetized with i.p. sodium phenobarbital, after which the common left sciatic nerve was exposed at the level of the middle of the thigh by blunt dissection through the biceps femoris and proximal to the sciatic trifurcation. Four ligatures (4.0 braided silk) were tied loosely around it with about 1mm spacing. The muscle was closed in layers and two wound clips were applied to close the skin incision. The wound was then covered with topical antibiotics.

Nociceptive testing

[0096] Static allodynia (a form of hyperlagesia) was measured using a method described by Chaplan *let al*, "Quantitative assessment of tactile allodinya in the rat paw", *J. Neurosci. Methods*, **53**, 55-63 (1994). A series of von Frey filaments of different buckling weight (i.e. the load required for the filament to bend) were applied to the plantar surface of the right hand paw. The starting filament had a buckling weight of 20g. Lifting of the paw was taken to be a positive result, in which case a filament with the next lowest buckling weight was used for the next measurement. The test was continued until a filament was found for which there was an absence of response for longer than 5 seconds whereas a re-test with the next heaviest filament gave a positive response. Animals were considered hyperalgesic if their thresholds were found to be <4g of those of comparable untreated rats, see Calcutt & Chaplan, "Spinal pharmacology of tactile allodynia in diabetic rats", *British J. Pharmacol*, **122**, 1478-1482 (1997).

Tissue Extraction

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[0097] STZ-treated, CCI-treated and control animals were anaesthetized with 4% halothane and perfused with ice-cold 0.9% saline containing 1% citric acid (pH adjusted to 7.4 with NaOH). The animals were decapitated and the lumbar spinal cord exposed. A 2-centimetre length of spinal cord ending at L6 (lumbar-6 forward) was removed from the spinal column. Attached dorsal root ganglia and contaminating spinal connective tissues were removed. The spinal cord tissue was snap frozen in dry ice and isopentane. In the experiments that follow, procedures on the streptozocintreated and control groups of animals are disclosed. It will be understood that for identification of CCI-treated animals the same experiments are performed, but using tissues from the CCI-treated animals and from control animals.

Total RNA Extraction

[0098] Total RNA was extracted from the pooled male rat tissues of the streptozocin-treated and control groups using the TRIZOL Reagent Kit (Life Technologies). Briefly, tissue samples were homogenised fully using a Polytron homogenizer in 1ml of TRIZOL reagent per 50-100mg of tissue. Homogenized samples were then incubated at room temperature for 5 minutes and phase separated using 0.2ml chloroform per 1ml TRIZOL reagent followed by centrifugation at 3,000g. The aqueous phase was transferred to a fresh tube and the RNA precipitated with an equal volume of isopropyl alcohol and followed by centrifugation at 10,000g. The RNA pellet was washed in 75% ethanol and re-centrifuged. The pellet was then air dried and re-suspended in water.

mRNA Extraction

[0099] In contrast to ribosomal RNA and transfer RNA, the vast majority of mRNAs of mammalian cells carry tracts of poly(A+) at their 3' termini. mRNAs can therefore be separated from the bulk of cellular RNA by affinity chromatography on oligo (dT)-cellulose. mRNA was extracted from Total RNA using the MESSAGEMAKER Kit (Life Technologies) in which mRNA (previously heated to 65°C in order to disrupt secondary structures and so expose the poly (A+) tails) was bound to oligo(dT) cellulose under high salt concentrations (0.5M NaCl) in a filter syringe. Unbound RNA was then washed away and the poly(A+) mRNA eluted in distilled water. A tenth of the volume of 7.5 M Ammonium Acetate, 50µg of glycogen/ml mRNA sample and 2 volumes of absolute alcohol were then added to the samples which were placed at -20°C overnight. Following precipitation, the mRNA was spun down at 12,000g for 30 minutes at 4°C. RNAase free water was used to re-suspend the pellets, which were then and stored at -80°C.

PCR SELECT

[0100] The technique used was based on that of the CLONTECH PCR Select Subtraction Kit. The following protocol was performed using STZ-treated lumbar spinal cord Poly A + RNA as the 'Tester' and Sham lumbar spinal cord poly A+ RNA as the 'Driver' (Forward Subtraction). A second subtraction experiment using the Sham lumbar spinal cord mRNA as the 'Tester' and STZ treated lumbar spinal cord mRNA as the 'Driver' (Reverse Subtraction) was performed in parallel using the same reagents and protocol. As a control for both experiments, the subtraction was also carried out using human skeletal muscle mRNA that had been provided by the manufacturer.

First-Strand cDNA Synthesis

[0101] $2 \mu g$ of PolyA⁺ RNA and $1 \mu l$ of cDNA synthesis primer (10 μM) were combined in a 0.5ml Eppendorf tube and sterile H₂O was added where necessary to achieve a final volume of 5 μl . The contents were mixed gently and incubated in a thermal cycler at 70°C for 2 min. The tubes were then cooled on ice for two minutes, after which 2 μl of

5X First-strand buffer, 1 μ I of dNTP mix (10 mM each), sterile H₂O and 1 μ I of AMV reverse transcriptase (20 units/ μ I) was also added. The tubes were then placed at 42°C for 1.5 hr in an air incubator. First-strand cDNA synthesis was terminated by placing the tubes on ice. (the human skeletal muscle cDNA produced by this step was used as the 'control driver' in later steps).

Second-Strand cDNA Synthesis

[0102] 48.4 μ l of Sterile H₂O, 16.0 μ l of 5X Second-strand buffer, 1.6 μ l of dNTP mix (10 mM) and 4.0 μ l of 20X Second-strand enzyme cocktail were added to each of the first-strand synthesis reaction tubes. The contents were then mixed and incubated at 16°C in a thermal cycler for 2 hr. 6 units (2 μ l) of T4 DNA polymerase was then introduced and the tubes were incubated for a further 30 min at 16°C. In order to terminate second-strand synthesis, 4 μ l of 20X EDTA/glycogen mix was added. A phenol:chloroform extraction was then carried out using the following protocol:[0103] 100 μ l of phenol:chloroform:isoamyl alcohol (25:24:1) was added to the tubes which were then vortexed thoroughly and centrifuged at 14,000 rpm for 10 min at room temperature. The top aqueous layer was removed and placed in a fresh tube. 100 μ l of chloroform:isoamyl alcohol (24:1) was then added to the aqueous layer and the tubes were again vortexed and centrifuged at 14,000 rpm for 10 min. 40 μ l of 4 M NH₄OAc and 300 μ l of 95% ethanol were then added and the tubes centrifuged at 14,000 rpm for 20 min. The supernatant was removed carefully, then 500 μ l of 80% ethanol was added to the pellet. The tubes were centrifuged at 14,000 rpm for 10 min and the supernatant was removed so that the pellet could be air-dried. The precipitate was then dissolved in 50 μ l of H₂O. 6 μ l was transferred to a fresh microcentrifuge tube. The remainder of the sample was stored at -20°C until needed.

Rsa I Digestion

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[0104] All products of the above procedures were subjected to a restriction digest, using the restriction endonuclease Rsa I, in order to generate ds cDNA fragments that are short and thus are optimal for subtraction hybridisation due to the standard kinetics of the hybridisation. Also, as Rsa I makes a double stranded cut in the middle of a recognition sequence, 'blunt ends' of a known nucleotide sequence are produced allowing ligation of adaptors onto these ends in a later step. The following reagents were added to the 6 μ l product of the second hybridisation (see above): 43.5 μ l of ds cDNA, 5.0 μ l 10X Rsa I restriction buffer and 1.5 μ l of Rsa I (10 units/ μ l). The reaction mixture was incubated at 37°C for 1.5 hr. 2.5 μ l of 20X EDTA/glycogen mix was used to terminate the reaction. A phenol:chloroform extraction was then performed as above (second-strand cDNA synthesis section). The pellet produced was then dissolved in 5.5 μ l of H₂O and stored at -20°C until needed. The preparation of the experimental 'Driver' cDNAs and the control skeletal muscle cDNA was thus completed.

Adaptor Ligation

[0105] The adaptors were not ligated to the driver cDNA.

[0106] 1 μl of each Rsa I-digested experimental cDNA (from the Rsa I Digestion above) was diluted with 5 μl of sterile water. Preparation of the control skeletal muscle tester cDNA was then undertaken by briefly centrifuging the tube containing control DNA (Hae III-digest of φX174 DNA [3 ng/ μl]) and diluting 2 μl of the DNA with 38 μl of sterile water (to 150 ng/ml). 1 μl of control skeletal muscle cDNA (from the Rsa I Digestion) was then mixed with 5 μl of the diluted φX174/ Hae III DNA (150 ng/ml) in order to produce the control skeletal muscle tester cDNA.

Preparation of the adaptor-ligated tester cDNA

[0107] A ligation master mix was prepared by combining 3 μ l of sterile water, 2 μ l of 5X ligation buffer and 1 μ l T4 DNA ligase (400 units/ μ l) per reaction. 2 μ l of adaptor 1 (10 μ M) was then added to 2 μ l of the diluted tester cDNA. To this, 6 μ l of the ligation master mix was also added. The tube was therefore labeled Tester 1-1. In a separate tube, 2 μ l of the adaptor 2R (10 μ M) was mixed with 2 μ l of the diluted tester cDNA and 6 μ l of the master mix. This tube was named Tester 1-2.

[0108] $2 \mu l$ of Tester 1-1 and $2 \mu l$ of Tester 1-2 were then placed into fresh tubes. These would later be used as the unsubtracted tester control. The remainder of the contents of Tester 1-1 and Tester 1-2 tubes were then centrifuged briefly and incubated at 16°C overnight. The ligation reaction was stopped by adding 1 μl of EDTA/glycogen mix and the samples were heated at 72°C for 5 min in order to inactivate the ligase. In doing so, preparation of the experimental and control skeletal muscle adaptor-ligated tester cDNAs was complete.

[0109] 1 μ I from each unsubtracted tester control was then removed and diluted into 1 ml of water. These samples were set aside as they were to be used later for PCR (see below). All of the samples were stored at -20°C

Analysis of Ligation efficiency

[0110] 1 μ I of each ligated cDNA was diluted into 200 μ I of water and the following reagents were then combined in four separate tubes:

Component	Tube:	1	2	3	4
Tester 1-1 (ligated to Adaptor 1)		1	1	•	-
Tester 1-2 (ligated to Adaptor 2R)		-	-	1	1
G3PDH 3' primer (10 µM)		1	1	1	1
G3PDH 5' primer (10 μM)		-	1	-	1
PCR primer 1 (10 μM)		1	-	1	-
Total volume μl		3	3	3	3

[0111] A master mix for all of the reaction tubes plus one additional tube was made up by adding 18.5 µl of sterile H₂O, 2.5 µl of 10X PCR reaction buffer, 0.5 µl of dNTP mix (10 mM), and 0.5 µl of 50X Advantage cDNA Polymerase Mix, per reaction, into a fresh tube. 22 µl of this master mix was then aliquotted into each of the 4 reaction tubes prepared above. The contents of the tubes were overlaid with 50 µl of mineral oil. The reaction mix was incubated in a thermal cycler at 75°C for 5 min in order to extend the adaptors. The following protocol was then carried out immediately in a thermal cycler (Perkin-Elmer GeneAmp PCR Systems 2400): 94°C for 30 sec (1 cycle), 94°C 10 sec, 65°C 30 sec and then 68°C 2.5 min (25 cycles)

First Hybridisation

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[0112] 1.5 µl of the Adaptor 1-ligated Tester 1-1 was combined with 1.5 µl of the Rsa I-digested driver cDNA, prepared earlier and 1 µl of 4X Hybridisation buffer. This process was then repeated combining the Adaptor 2R-ligated Tester 1-2 with the Rsa I-digested driver cDNA and 4X hybridisation buffer. The samples were incubated in a thermal cycler at 98°C for 1.5 min followed by incubation at 68°C for 8 hr.

Second Hybridisation

[0113] 1 μ I of Driver cDNA (i.e. the *Rsa* I-digested cDNA (see above)), 1 μ I 4X Hybridisation buffer and 2 μ I Sterile H₂O were all combined in a fresh tube. 1 μ I of this mix was then removed and placed in a new tube, overlaid with 1 drop of mineral oil and incubated at 98°C for 1.5 min in order to denature the driver. The following procedure was used to simultaneously mix the driver with hybridisation samples 1 and 2 (prepared in the first hybridisation), thus ensuring that the two hybridisation samples were mixed together only in the presence of freshly denatured driver: A micropipettor was set at 15 μ I. The pipette tip was then touched onto the mineral oil/sample interface of the tube containing hybridisation sample 2. The entire sample was drawn partway into the tip before it was removed from the tube in order to draw a small amount of air into the tip. The pipette tip was then touched onto the interface of the tube containing the freshly denatured driver (i.e. the tip contained both samples separated by a small pocket of air) before the entire mixture was transferred to the tube containing hybridisation sample 1. The reaction was then incubated at 68°C overnight. 200 μ I of dilution buffer was added to the tube, which was then heated in a thermal cycler at 68°C for 7 min. The product of this second hybridisation was stored at -20°C.

PCR Amplification

[0114] Seven PCR reactions were set up: (1) The forward-subtracted experimental cDNA, (2) the unsubtracted tester control (see preparation of the adaptor ligated tester cDNA), (3) the reverse-subtracted experimental cDNA, (4) the unsubtracted tester control for the reverse subtraction, (5) the subtracted control skeletal muscle cDNA, (6) the unsubtracted tester control for the control subtraction, and (7) the PCR control subtracted cDNA (provided in the kit). The PCR control subtracted cDNA was required to provide a positive PCR control as it contained a successfully subtracted mixture of *Hae* III-digested \$\phi\$X174 DNA.

[0115] The PCR templates were prepared by aliquotting 1 μ l of each diluted cDNA (i.e., each subtracted sample from the second hybridisation and the corresponding diluted unsubtracted tester control produced by the adaptor ligation see above) into an appropriately labeled tube. 1 μ l of the PCR control subtracted cDNA was placed into a fresh tube. A master mix for all of the primary PCR tubes, plus one additional reaction, was then prepared by combining 19.5 μ l of sterile water, 2.5 μ l of 10X PCR reaction buffer, 0.5 μ l of dNTP Mix (10 mM), 1.0 μ l of PCR primer 1 (10 μ M)

and 0.5 µl of 50X Advantage cDNA Polymerase Mix. 24 µl of Master Mix was then aliquotted into each of the 7 reaction tubes prepared above and the mixture was overlaid with 50 µl of mineral oil, before being incubated in a thermal cycler at 75°C for 5 min in order to extend the adaptors. Thermal cycling was then immediately started using the following protocol: 94°C 25 sec (1 cycle), 94°C 10 sec, 66°C 30 sec and 72°C 1.5 min (32 cycles).

[0116] 3 μ l of each primary PCR mixture was then diluted in 27 μ l of H₂O, 1 μ l of each of these dilutions was then placed into a fresh tube.

[0117] A master mix for the secondary PCRs, (plus an additional reaction) was set up by combining 18.5 μ l of sterile water, 2.5 μ l of 10X PCR reaction buffer, 1.0 μ l of Nested PCR primer 1 (10 μ M), 1.0 μ l of Nested PCR primer 2R (10 μ M), 0.5 μ l of dNTP Mix (10 mM) and 0.5 μ l of 50X Advantage cDNA Polymerase Mix per reaction. 24 μ l of this Master Mix was then added into each reaction tube containing the 1 μ l diluted primary PCR mixture. The following PCR protocol was then carried out: 94°C 10 sec, 68°C 30 sec and 72°C 1.5 min (12 cycles). The reaction products were then stored at -20°C.

Ligation into a Vector/Transformation & PCR

[0118] The products of the PCR amplification (enriched for differentially expressed cDNAs) were ligated into the pCR2.1-TOPO vector using a T/A cloning kit (Invitrogen), transformed into TOPO One Shot competent cells according to the manufacturers protocol and grown up on LB (Luria-Bertani) Agar plates overnight at 37°C. 1,000 colonies were then individually picked (using fresh sterile tips) and dipped into 5 μ l of sterile water which had been aliquotted previously into 96 well PCR plates. The water/colonies were heated in a thermal cycler at 100°C for 10 minutes in order to burst the cells, thus releasing the plasmids containing a differentially expressed cDNA insert. The 5 μ l of water/plasmid was then used as a template in a PCR reaction (see below) using M13 Forward and Reverse primers (10 ng/ μ l), complementary to the M13 site present on either side of the cloning site on the vector. 5 μ l of the PCR product was then run on a 2% agarose gel and stained by ethidium bromide. PCR products of an amplified insert were identified and 5 μ l of the remainder of the PCR product (i.e. from the 15 μ l that had not been run on the gel) was diluted 1/10 with water. 5 μ l of the diluted PCR product was then used as a template in a sequencing reaction.

Sequencing

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[0119] A sequencing reaction containing M13 primer (3.2 pmol/µl), 'BigDye' reaction mix (i.e. AmpliTaq® DNA polymerase, MgCl₂, buffer and fluorescent dNTPs [each of the four deoxynucleoside triphosphates is linked to a specific fluorescent donor dye which in turn is attached to a specific acceptor dye]) and cDNA template (diluted PCR product) was set up. The reaction was carried out on a thermal cycler for 25 cycles of 10 seconds at 96°C, 20 seconds at 50°C and 4 minutes at 60°C. Each reaction product was then purified through a hydrated Centri-Sep column, and lyophilised. The pellets were resuspended in Template Supression Reagent and sequenced on an ABI Prism 310 Genetic Analyser. The analyser uses an ion laser to excite the specific donor dye that transfers its energy to the acceptor dye, which emits a specific energy spectrum that can be read by the sequencer.

[0120] The differentially expressed genes of the streptozocin-induced diabetes experiment and of the CCI experiment were sequenced at Parke-Davis, Cambridge and at the applicant's core sequencing facility in Ann Arbor, MI, USA.

Bioinformatics

[0121] The sequencing results were analysed using the computer program CHROMAS in which the vector and adaptor sequences were clipped off, leaving only the nucleotide sequence of the differentially expressed gene. Each sequence was then checked for homology to known genes, Expressed Sequence Tags (ESTs) and Proteins using various Basic Local Alignment Search Tool (BLAST) searches against the Genbank sequence database at the National Centre for Biotechnology Information, Bethesda, Maryland, USA (NCBI).

[0122] Lists were derived called STZup and STZdown and CClup and CCldown that contain the nucleic acid sequences from the forward and back subtracted libraries respectively. In each list there are given accession numbers and descriptions for the known rat genes identified, and where available corresponding mouse or human genes. Sequences that are considered to be of interest and that are down-regulated both in a streptozocin-induced diabetes model and in a chronic constrictive injury pain model are identified in Tables I - VI above and are listed below.

[0123] References have been given where available for the sequences that have been found, and sequence listings have been given in the form in which they currently appear in publicly searchable databases e.g. the NCBI databases (National Center for Biotechnology Information, National Library of Medicine, National Institutes of Health, Bethesda, Maryland, MD 20894, USA, www.ncbi.nlm.nih.gov). These sequence listings are given for the purposes of identification only. The invention includes the use of subsequently revised versions of the above sequences (which may incorporate small differences to the version set out herein) and homologous sequences or similar proteins in other species as

determined by a high percentage identity (e.g. above 50%, preferably above 90%), length of alignment and functional equivalence.

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SEQUENCE LISTING

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 -	Ala Thr Lys Leu Gly Pro	Val Gly Gly Val Phe Asn	Leu Ala Met Val
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	Val Asn Lys Pro Lys Tyr	Asn Gly Thr Leu Asn Leu	Asp Arg Ala Thr
	1985 1990	1995	2000
55	Arg Glu Ala Cys Pro Glu	Leu Asp Tyr Phe Val Ala	Phe Ser Ser Val
	2005	2010	2015

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A65	Gly Leu Ala Ala Gly Asn Ser Gly Gly Pro Ala Gly Thr Thr Ala Pro 485	10	450 455 460
Ser Thr Ala Pro Gly Glu Asp Thr Asn Pro Gln Gly Gly Ala Ala Glu 510 Pro Gly His Gln Gln Phe IIe Gln Gln Met Leu Gln Ala Leu Ala Gly 515 Val Asn Pro Gln Leu Gln Ser Pro Glu Val Arg Phe Gln Gln Gln Leu Gln 530 Z5 Glu Gln Leu Ser Ala Met Gly Phe Leu Lys Arg Glu Pro Asn Leu Gln 545 555 556 Ala Leu Ile Ala Thr Gly Gly Asp Ile Asn Ala Ala Ile Glu Arg Leu 565 Leu Gly Ser Gln Pro Ser 580 Leu Gly Ser Gln Pro Ser 580 220 <2210> 101 2211> 1843 2212> DNA 2213> Mus musculus 43 Leu Caaatctca tattgaccaa dectagaagg teacggtgaa gagcgctaaa gagcgcctaaa gagcgcctaaa gagcaaggaa thaacatgacg teacaggtgaa aacaaatac cacaggc teacaggtgaa gattaacaaggc gaagaaggaa teacaggtgaa gacttaagg teacaggtgaa cacagttca daaacaggcc gaagagtaa teacgagaa aacaaatac cacaggc teacaggtgaa agttgaccaa gagtcacag ctcttgaaga tacaccaagg tetacacagaa teacaggtgaa caccaagaa teacaggtgaa caccaagaa teacaggtgaa caccaagaa teacaggtgaa aacaaatac cacaggac tectgaaca 360 gttaacaagg tectctgaaag aacaccaaga teacaggtgaa caccaagaa teacaggtgaa aacaaatac caccagg teacaggaaca agttaacaaacagc gtaacaaaggc tectgaagaa aacaaatac caccaggac tectgaaca 360 gttaacaaagg tectctgaaaga aacaaatac caccaagaac tecttgagaaca agttaacaaacac 300 aaaacaaggcc gaagaacaaat teagaccaaga tetagagaac tecttgaaca 360 gttaacaaa cecttgtgaagaa caccaacaga tectagaaaa accaacattc 480 teatggaaca cecttgagaacaac cecacaggt gaagaagaagaa attaagacaagac tectagaaca cecacagagac taccaagagaa agcagcaaa agcagaacaa cecacacattc 480 teatggacaa cecacacacatac cacaagagacaac cecacaagagacaac cacaagagacaacaacaacaagacaagacaacaacacacaaca	Ser Thr Ala Pro Gly Glu Asp Thr Asn Pro Gly Gly Ala Ala Glu 510		= "
Pro Gly His Gln Gln Phe Ile Gln Gln Met Leu Gln Ala Leu Ala Gly 515 520	Pro Gly His Gln Gln Phe Ile Gln Gln Met Leu Gln Ala Leu Ala Gly 515 Val Asn Pro Gln Leu Gln Ser Pro Glu Val Arg Phe Gln Gln Gln Leu 530 Z5 Glu Gln Leu Ser Ala Met Gly Phe Leu Lys Arg Glu Pro Asn Leu Gln 540 Ala Leu Ile Ala Thr Gly Gly Asp Ile Asn Ala Ala Ile Glu Arg Leu 565 Ala Leu Gly Ser Gln Pro Ser 555 Leu Gly Ser Gln Pro Ser 580 Z20> Z20> Z20> Z20> Z210> 101 C211> 1843 Z212> DNA Z213> Mus musculus Z20> 15		
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Claims

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1. Use of:

- (a) an isolated gene sequence that is down-regulated in the spinal cord of a mammal in response to mechanistically distinct first and second models of neuropathic or central sensitization pain;
 - (b) an isolated gene sequence comprising a nucleic acid sequence of any of Tables I to VI;
 - (c) an isolated gene sequence having at least 80% sequence identity with a nucleic acid sequence of any of Tables I to VI:
 - (d) an isolated nucleic acid sequence that is hybridizable to any of the gene sequences according to (a), (b) or (c) under stringent hybridisation conditions;
 - (e) a recombinant vector comprising a gene sequence or nucleic acid sequence according to any one of (a) to (d);
 - (f) a host cell containing the vector according to (e);
- (g) a non-human animal having in its genome an introduced gene sequence or nucleic acid sequence or a removed or down-regulated gene sequence or nucleic acid sequence according to any one of (a) to (d);
 - (h) an isolated polypeptide comprising an amino acid sequence at least 90% identical to an amino acid sequence encoded by a nucleotide sequence according to any one of (a) to (d), or a polypeptide variant thereof

with sequential amino acid deletions from the C terminus and/or the N-terminus;

- (i) an isolated polypeptide encoded by a nucleotide sequence according to any one of (a) to (d); or
- (i) an isolated antibody that binds specifically to a polypeptide according to (h) or (i);
- 5 in the screening of compounds for the treatment of pain, or for the diagnosis of pain.
 - 2. Use according to claim 1, wherein the isolated gene sequence is down-regulated both in response to streptozocininduced diabetes and in response to surgical injury of a nerve leading to the spine.
- 10 3. Use according to claim 1 or 2 wherein the isolated gene sequence encodes a kinase.
 - 4. Use according to claim 1, 2 or 3, wherein the isolated gene sequence encodes an expression product or fragment thereof of pyruvate kinase, M1 and M2 subunits (M24359; X97047; X56494).
- Use according to claim 1 or 2, wherein the isolated gene sequence encodes an expression product or fragment thereof of a receptor.
 - 6. Use according to claim 1, 2 or 5, wherein the isolated gene sequence encodes dopamine receptor D.sub.1 (I58000).
- 20 7. Use according to claim 1 or 2, wherein the isolated gene sequence encodes a transporter.
 - Use according to claim 1, 2 or 7, wherein the isolated gene sequence encodes differentiation-associated Natdependent inorganic phosphate cotransporter (AF271235) or putative vacuolar assembly protein VSP41 gene (U87309).
 - 9. Use according to claim 1 or 2, wherein the isolated gene sequence encodes a G-protein coupled receptor protein.
 - Use according to claim 1, 2 or 9, wherein the isolated gene sequence encodes Git1 (G-protein-coupled receptor kinase-interactor 1; GPCR kinase-associated ADP-ribosylation factor) (AF085693).
 - 11. Use according to claim 1 or 2, wherein the isolated gene sequence encodes a DNA-binding protein.
 - Use according to claim 1, 2 or 11, wherein the isolated gene sequence encodes putative histone H3.3A (X91866; M11354).
 - 13. Use according to claim 1 or 2, wherein the isolated gene sequence encodes a ligase.
 - 14. Use according to claim 1, 2 or 13, wherein the isolated gene sequence encodes 3-Hydroxy 3-methylglutaryl coenzyme A synthase, cytosolic (X52625), acyl-CoA synthetase II, brain (D360666) famesyl diphosphate synthase (M34477), bendless protein (AB032739; E12457), fatty acid synthase (X62888), glutamine synthetase (EC 6.3.1.2) (M91652), or putative seryl-tRNA synthetase (X91257).
 - 15. Use according to claim 1 or 2, wherein the isolated gene sequence encodes a lyase.
- 45 16. Use according to claim 1, 2 or 15, wherein the lyase is enolase, alpha alpha, non-neuronal (NNE) (X02610; X52379; M14328).
 - 17. Use according to claim 1 or 2, wherein the isolated gene sequence encodes an oxidoreductase.
- 18. Use according to claim 1, 2 or 17, wherein the isolated gene sequence encodes aldose reductase, lens (AREC 11.1.21) (X05884) cytochrome-c oxidase I, mitochondrial (S79304), lactate dehydrogenase-B (LDH-B) (U07181; X51905; Y00711), putative cytochrome c oxidase VIB (EC 1.9.3.1) (X13923), putative NADH: ubiquinone oxidore-ductase PGIV subunit (AF044953), putative succinate dehydrogenase flavoprotein (AF095938; AF171022), putative ubiquinol—cytochrome-c reductase (EC 1.10.2.2) core protein II (J04973) or stearoyl-coA desaturase 2 (AB032243; M26270).
 - 19. Use according to claim 1 or 2, wherein the isolated sequence encodes a transferase.

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- 20. Use according to claim 1, 2 or 19, wherein the isolated sequence encodes ribophorin I (X05300) or sulfotransferase-like protein (AF188699).
- 21. Use according to claim 1 or 2, wherein the isolated sequence encodes a hydrolase.
- 22. Use according to claim 1, 2 or 21, wherein the isolated sequence encodes ATP synthase, H+, alpha subunit, mitochondrial (EC 3.6.1.34) (X56133), F1F0 ATPase delta subunit (U00926), putative dihydropyrimidinase related protein (D78013), heat shock protein 90 (S45392; M18186; M16660), or putative ribonuclease III (AF116910).
- 10 23. Use according to claim 1 or 2, wherein the isolated sequence encodes myelin basic protein S (MBP S) (K00512), transferring (D38380), neurofilament, light molecular weight (NF-L) (AF031880), myelin-associated glycoprotein (MAG) (M16800; M31811), NF-M middle molecular weight neurofilament protein (M18628) neuro-degeneration associated- protein 1 (D32249), S-100 protein β-subunit (X01090), microtubule-associated protein 1b (Map 1b) (X60370; L06237), putative cdc 37 homolog (D26564), putative ras-related protein Rab-5c (U11293), putative 15 gelsolin (J04953), Cd81 antigen (target of antiproliferative antibody 1) (U19894; X59047; M33680), Mobp81 (Myelin-associated/Oligo-dendrocytic basic protein 81) (X87900), syntaxin binding protein n-secl, sec1 homolog, Ainternexin (M73049), putative β-sarcoglycan A3b (AB024921), CGI-78 protein (AF151835), KIAA0143 (D63477), septin 2 (D50918), Nucleobindin (Z36277), myelin protein SR13 (M69139; S55427), B-Actin, cytoplasmic (V01217; X03672), ly6/neurotoxin (Lynx1) homolog (AD141377), astrocytic phosphoprotein; PFA 15 gene (AJ243949; 20 X86694), PLIC-1 (AF177345), Nfx1 (tip associating protein (TAP) gene) (AF093139; AF093140), A-Crystallin B (U04320; M73741; M28638), heat shock-like protein 70 kD (X70065; U73744; Y00371), tau microtubule-associated protein (X79321), myelin, Schwann cell, Perioheral (P-0) (K03242), B-Tubulin class 1 (AB011679; X04663; AF14139), putative long-chain polyunsaturated fatty acid elongation enzyme (Helo1) or peptidylglycine alphaamidating monooxygenase 1.
 - 24. A non-human animal having in its genome an introduced gene sequence or a removed or down-regulated gene sequence, said sequence being down-regulated in the spinal cord of a mammal in response to first and second models of neuropathic or central sensitisation pain.
- 25. A non-human animal according to claim 24, wherein said gene sequence becomes down regulated both in response to streptozocin induced diabetes and in response to chronic constriction injury.
 - 26. A non-human animal according to claim 24 or 25, wherein the introduced gene sequence is according to any of claims 1 to 23.
 - 27. A non-human animal according to any one of claims 24 to 26 which is C. elegans.
 - 28. A kit comprising;

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- (a) affinity peptide and/or ligand and/or substrate for an expression product of a gene sequence that is down-regulated in the spinal cord of a mammal in response to a mechanistically distinct first and second models of neuropathic or central sensitization pain; and
- (b) a defined quantity of an expression product of a gene sequence that is down-regulated in the spinal cord of a mammal both in response to first and second models of neuropathic or central sensitization pain, for simultaneous, separate or sequential use in detecting and/or quantifying down-regulation of a gene sequence in the spinal cord of a mammal in response to first and second models of neuropathic or central sensitization pain.
- 29. A kit according to claim 28, wherein the gene sequence is defined in any one of claims 1 to 23.
 - 30. A kit comprising:
 - (a) nucleic acid sequences capable of hybridization to a nucleic acid sequence that is down-regulated in the spinal cord of a mammal in response to first and second models of neuropathic or central sensitization pain; and (b) a defined quantity of one or more nucleic acid sequences capable of hybridization to a nucleic acid sequence that is down-regulated in the spinal cord of a mammal in response to first and second models of neuropathic or central sensitization pain,

for simultaneous, separate or sequential use in detecting and/or quantifying down-regulation of a gene sequence in the spinal cord of a mammal in response to first and second models of neuropathic or central sensitization pain.

- 5 31. The kit of claim 30, wherein the gene sequence is according to any of claims 1 to 23.
 - 32. A compound that modulates the action of an expression product of a gene sequence that is down-regulated in the spinal cord of a mammal in response to first and second models of neuropathic or central sensitization pain.
- 33. A compound according to claim 32 wherein the gene sequence is listed in Tables I to VI.
 - 34. A compound according to claim 32 or 33 wherein the nucleotide sequence is according to any one of claims 1 to 23.
 - 35. A compound according to any one of claims 32 to 34 for use as a medicament.

36. A compound according to any one of claims 32 to 35 for the treatment or diagnosis of pain.

- 37. A pharmaceutical composition comprising a compound according to any one of claims 32 to 36 and a pharmaceutically acceptable carrier or diluent.
- **38.** Use of a compound according to any one of claims 32 to 36 in the manufacture of a medicament for the treatment or diagnosis of pain.
- 39. Use of a compound according to any one of claims 32 to 36 in the manufacture of a medicament for the treatment or diagnosis of chronic pain.
 - **40.** A method of treatment of pain, which comprises administering to a patient an effective amount of a compound according to any one of claims 32 to 36.

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(11) EP 1 284 298 A3

(12)

EUROPEAN PATENT APPLICATION

(88) Date of publication A3: 26.05.2004 Bulletin 2004/22

(51) Int CI.7: **C12Q 1/68**, C07K 14/47, C12N 5/10, C12N 15/85

(43) Date of publication A2: 19.02.2003 Bulletin 2003/08

(21) Application number: 02255229.3

(22) Date of filing: 26.07.2002

(84) Designated Contracting States:

AT BE BG CH CY CZ DE DK EE ES FI FR GB GR
IE IT LI LU MC NL PT SE SK TR
Designated Extension States:

AL LT LV MK RO SI

(30) Priority: 27.07.2001 GB 0118354 07.02.2002 GB 0202883

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(54) Identification and use of molecules implicated in pain

(57) The invention relates to the use of:

(a) an isolated gene sequence that is down-regulated in the spinal cord of a mammal in response to mechanistically distinct first and second models of neuropathic or central sensitization pain;

(b) an isolated gene sequence comprising a nucleic acid sequence of any of Tables I to VI;

(c) an isolated gene sequence having at least 80% sequence identity with a nucleic acid sequence of any of Tables I to VI;

(d) an isolated nucleic acid sequence that is hybridizable to any of the gene sequences according to (a), (b) or (c) under stringent hybridisation conditions:

(e) a recombinant vector comprising a gene sequence or nucleic acid sequence according to any one of (a) to (d);

(f) a host cell containing the vector according to (e);
(g) a non-human animal having in its genome an introduced gene sequence or nucleic acid sequence or a removed or down-regulated gene sequence or nucleic acid sequence according to any one of (a) to (d);

(h) an isolated polypeptide comprising an amino acid sequence at least 90% identical to an amino acid sequence encoded by a nucleotide sequence ac-

cording to any one of (a) to (d), or a variant polypeptide thereof with sequential amino acid deletions from the C terminus and/or the N-terminus;

(i) an isolated polypeptide encoded by a nucleotide sequence according to any one of (a) to (d); or

(j) an isolated antibody that binds specifically to a polypeptide according to (h) or (i);

in the screening of compounds for the treatment of pain, or for the diagnosis of pain.

The invention also relates to the use of naturally occurring compounds such as peptide ligands of the expression products of the above gene sequences and their associated signal transduction pathways for use in the treatment of pain.



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which under Rule 45 of the European Patent Convention EP 62 25 5229 shall be considered, for the purposes of subsequent proceedings, as the European search report

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	DOCUMENTS CONSIDI	ERED TO BE RELEVANT		
Category	Citation of document with in of relevant passa	dication, where appropriate, ges	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int.Cl.7)
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¥		ve action and a cord gene expression experimental diabetes" ch 1998 (1998-03), 62337	1-3, 24-31	TECHNICAL FIELDS SEARCHED (INLCI.7) C12Q C07K C12N
The Sear not comp be carried Claims se Claims se Claims no		application, or one or more of its claims, does a meaningful search into the state of the art c y, for these claims.		
	Place of search	Date of completion of the search		Examiner
	Munich	3 December 2003	Hei	rmann, P
X : par Y : par doc A : tecl O : nor	ATEGORY OF CITED DOCUMENTS ticularly relevant if taken alone ticularly relevant if combined with anotument of the same category inhological background bewritten disclosure immediate document	T: theory or principl E: earlier patent do after the filing da her D: document cited L: document cited	le underlying the cument, but publication in the application or other reasons	invention ished on, or

PO FORM 1503 ICI 82 (P04C)



INCOMPLETE SEARCH SHEET C

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The following comments apply to the invention for which a search has been performed - i.e. claims 1, 2, 24-40 (in part) and 3-4 (in full) relating to the first invention of the application.

The applicant should therefore bear in mind that, should he pay additional search fees, further objections leading to incomplete search might as well arise during the search phase of the further inventions.

Claim(s) searched completely: 2-4, 25

Claim(s) searched incompletely: 1, 24, 25, 27-31

Claim(s) not searched: 32-40

Reason for the limitation of the search:

a) Article 52 (4) EPC - Method for treatment of the human or animal body by therapy (claim 40).

Although claim 40 is directed to a method of treatment of the human/animal body (Article 52(4) EPC), the search could have been been carried out and based on the alleged effects of the compound/composition. However, in view of the lack of clarity of said claim (see item (f) hereinbelow) no search has been carried out.

- b) The isolated gene sequence of part (a) of claim 1 is only defined by a result achieved in special conditions, and the description is silent as to any other characteristic features pertaining to said isolated gene sequence. Claim 1 therefore lacks clarity (Article 84 EPC), and could only be searched as it relates to the isolated gene sequences of table I encoding a kinase, and derivative thereof as listed in parts (c)-(j) of claim 1, all relating to kinase encoding nucleic acid sequences.
- c) The non-human animal of present claims 24 or 25 or the C. elegans of claim 27 is defined by reference to the following parameter:
- the modification of the presence or expression of a gene sequence which is downregulated in the spinal cord of a mammal in response to two models of neuropathic pain and claim 25, as for it, limits those two models to the streptozocin induced-diabetes and the chronic constriction injury.

The use of that parameter in the present context is considered to lead to a lack of clarity within the meaning of Article 84 EPC (see also Guidelines C-III, 4.7a). For the same reasons as those hereinabove under item (b), the search for claims 24, 25 and 27 has been restricted to



INCOMPLETE SEARCH SHEET C

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non-human animal wherein the introduced gene sequence is one of the sequences listed in Table I, encoding a kinase.

- d) The expression product of a gene sequence and the nucleic acid sequence of the kits of claims 28-31 are only defined by a parameter i.e. the expression product of said gene sequence or the nucleic acid sequence that is downregulated in the spinal cord of a mammal in response to two models of neuropathic pain. The use of that parameter in the present context is considered to lead to a lack of clarity within the meaning of Article 84 EPC (see also Guidelines C-III, 4.7a). For the same reasons as those hereinabove under item (b), the search for claims 28-31 has been restricted respectively to the product of the gene sequences given in Table I and to the sequences given in Table I themselves.
- e) Moreover in independent claim 28 none of the characteristic features of the "affinity peptide", "ligand" and "substrate" for the expression product are given in said claim, and the description is also silent as to said eventual characteristics. Therefore the skilled person would not clearly understand which compounds are falling under the scope of said claim. Said claim therefore lacks clarity (Article 84 EPC) to such an extent that a meaningful search on its entire scope could not be performed. Said search has been restricted to substrates known for the kinases encoded by the gene sequences listed in table I.
- f) Claims 32-36 relate to compounds only defined by the result to be achieved and the claims as well as the description do not relate to any features which would allow the skilled person to clearly understand which compounds are falling under the scope of said claims. Said claims therefore lack clarity (Article 84 EPC) to such an extent that no meaningful search could be performed.

As claims 37-40 are depending upon one or more of claims 32-36, said claims therefore lack clarity (Article B4 EPC) and could not be searched either.



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	DOCUMENTS CONSIDERED TO BE RELEVANT	NSIDERED TO BE RELEVANT CLASSIFICATION (Internal Control of Character)			
Category	Citation of document with indication, where appropriate, of relevant passages	Relevant to claim	- · · · · · · · · · · · · · · · · · · ·		
Y	BLACKBURN-MUNRO G ET AL: "The sodium channel auxiliary subunits betal and beta2 are differentially expressed in the spinal cord of neuropathic rats" NEUROSCIENCE, vol. 90, no. 1, April 1999 (1999-04), pages 153-164, XP002262713 ISSN: 0306-4522 * the whole document *	1-3, 24-31			
A	NAKAMURA JIRO ET AL: "A protein kinase C-beta-selective inhibitor ameliorates neural dysfunction in streptozotocin-induced diabetic rats" DIABETES, vol. 48, no. 10, October 1999 (1999-10), pages 2090-2095, XP002260473 ISSN: 0012-1797 * the whole document *	1-3, 24-31	TECHNICAL FIELDS SEARCHED (Int.Cl.7)		
A	LUO Z DAVID ET AL: "Neuronal nitric oxide synthase mRNA upregulation in rat sensory neurons after spinal nerve ligation: Lack of a role in allodynia development" JOURNAL OF NEUROSCIENCE, vol. 19, no. 21, 1 November 1999 (1999-11-01), pages 9201-9208, XP002262714 ISSN: 0270-6474 * the whole document *	1-3, 24-31			



Application Number

EP 02 25 5229

	DOCUMENTS CONSIDERED TO BE RELEVANT	CLASSIFICATION OF THE APPLICATION (Int.Cl.7)			
Category	 Citation of document with indication, where appropriate, of relevant passages 	Relevant to daim			
A	FOX ALYSON ET AL: "Critical evaluation of the streptozotocin model of painful diabetic neuropathy in the rat" PAIN, vol. 81, no. 3, June 1999 (1999-06), pages 307-316, XP002260439 ISSN: 0304-3959 * the whole document *	1-3, 24-31			
A	COURTEIX C ET AL: "Streptozocin-induced diabetic rats: Behavioural evidence for a model of chronic pain" PAIN, ELSEVIER SCIENCE PUBLISHERS, AMSTERDAM, NL, vol. 53, no. 1, 1993, pages 81-88, XP009020332 ISSN: 0304-3959 * the whole document *	1-3, 24-31	TECHNICAL FIELDS SEARCHED (Int.Cl.7)		
A	MALMBERG ANNIKA B ET AL: "Preserved acute pain and reduced neuropathic pain in mice lacking PKC-gamma" SCIENCE (WASHINGTON D C), vol. 278, no. 5336, 1997, pages 279-283, XP002260438 ISSN: 0036-8075 * the whole document *	1-3, 24-31			
A	WO 00/63427 A (CUVILLIER GWLADYS ;DEVGEN NV (BE); BOGAERT THIERRY (BE); PLATTEEUW) 26 October 2000 (2000-10-26) * the whole document *	1-3, 24-31			
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TECHNICAL FIELDS SEARCHED (Int.Cl.7)
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CLAIMS INCURRING FEES
The present European patent application comprised at the time of filing more than ten claims.
Only part of the claims have been paid within the prescribed time limit. The present European search report has been drawn up for the first ten claims and for those claims for which claims fees have been paid, namely claim(s):
No claims fees have been paid within the prescribed time limit. The present European search report has been drawn up for the first ten claims.
LACK OF UNITY OF INVENTION
The Search Division considers that the present European patent application does not comply with the requirements of unity of invention and relates to several inventions or groups of inventions, namely:
see sheet B
All further search fees have been paid within the fixed time limit. The present European search report has been drawn up for all claims.
As all searchable claims could be searched without effort justifying an additional fee, the Search Division did not Invite payment of any additional fee.
Only part of the further search fees have been paid within the fixed time limit. The present European search report has been drawn up for those parts of the European patent application which relate to the inventions in respect of which search fees have been paid, namely claims:
None of the further search fees have been paid within the fixed time limit. The present European search report has been drawn up for those parts of the European patent application which relate to the invention first mentioned in the claims, namely claims: 1, 2, 24-40 (in part) 3-4 (in full)



LACK OF UNITY OF INVENTION SHEET B

Application Number

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The Search Division considers that the present European patent application does not comply with the requirements of unity of invention and relates to several inventions or groups of inventions, namely:

Invention 1: 1, 2, 24-40 (in part) 3-4 (in full)

Invention 1 relates to: i) the use of an isolated gene sequence encoding a kinase according to Table I, or derivative thereof such as recombinant vector, host cells or non-human animal comprising said isolated gene sequence, or isolated polypeptide encoded by, or comprising an isolated polypeptide encoded by said isolated gene sequence or isolated antibody directed against the above cited polypeptides, in the screening of compounds for the treatment of pain, or for the diagnosis of pain; ii) a non-human animal having introduced in its genome, said isolated gene sequence; iii) kits comprising either the expression product of said isolated gene sequence and its substrate, ligand or affinity peptide; iv) a compound that modulates the action of the expression product of said selected gene sequence; v) a pharmaceutical composition comprising said compound; vi) the use of said compound for the treatment or the diagnosis of pain; and vii) a method of treatment of pain which comprises administering to a patient an effective amount of said compound.

Invention 2: 1, 2, 24-40 (in part) 5-6 (in full)

Invention 2 is equivalent to invention 1 for an isolated gene sequence encoding an expression product or a fragment thereof of a receptor according to Table II.

Invention 3: 1, 2, 24-40 (in part) 7-8 (in full)

Invention 3 is equivalent to invention 1 for an isolated gene sequence encoding a transporter according to Table III.

Invention 4: 1, 2, 24-40 (in part) 9-10 (in full)

Invention 4 is equivalent to invention 1 for an isolated gene sequence encoding a G-protein coupled receptor protein according to Table IV.

Invention 5: 1, 2, 24-40 (in part) 11-12 (in full)

Invention 5 is equivalent to invention 1 for an isolated gene sequence encoding a DNA-binding protein according to Table V.

Invention 6: 1, 2, 24-40 (in part) 13-14 (in full)



LACK OF UNITY OF INVENTION SHEET B

Application Number

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The Search Division considers that the present European patent application does not comply with the requirements of unity of invention and relates to several inventions or groups of inventions, namely:

Invention 6 is equivalent to invention 1 for an isolated gene sequence encoding a ligase according to rows 1-7 of Table VI.

Invention 7: 1, 2, 24-40 (in part) 15-16 (in full)

Invention 7 is equivalent to invention 1 for an isolated gene sequence encoding a lyase according to row 8 of Table VI.

Invention 8: 1, 2, 24-40 (in part) 17-18 (in full)

Invention 8 is equivalent to invention 1 for an isolated gene sequence encoding an oxido-reductase according to rows 9-16 of Table VI.

Invention 9: 1, 2, 24-40 (in part) 19-20 (in full)

Invention 9 is equivalent to invention 1 for an isolated gene sequence encoding a transferase according to rows 17-18 of Table VI.

Invention 10: 1, 2, 24-40 (in part) 21-22 (in full)

Invention 10 is equivalent to invention 1 for an isolated gene sequence encoding a hydrolase according to rows 19-21 and 54 of Table VI.

Invention 11: 1, 2, 23-40 (in part)

Invention 11 is equivalent to invention 1 for an isolated gene sequence encoding a protein according to 23rd row of Table VI.

Inventions 12-41: 1, 2, 23-40 (in part)

Inventions 12-41 are equivalent to invention 1 for an isolated gene sequences encoding a protein according to the 24th-53rd rows of Table VI respectively.

ANNEX TO THE EUROPEAN SEARCH REPORT ON EUROPEAN PATENT APPLICATION NO.

EP 02 25 5229

This annex lists the patent family members relating to the patent documents cited in the above-mentioned European search report. The members are as contained in the European Patent Office EDP file on The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

03-12-2003

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